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54) Title: TRUNCATED VEGF-RELATED	PROTEINS					
		VEGF-B				
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The present invention provides novel truncated forms of vascular endothelial growth factor–related proteins (VRPs or VRPs) which are useful for the stimulation of angiogenesis *in vitro* and *in vivo*. The invention also provides nucleic acids encoding such novel truncated VRPs and methods of producing truncated VRPs. Pharmaceutical compositions comprising truncated VRPs and methods of gene therapy using the nucleic acids which code for truncated VRPs may be useful for the treatment of heart disease and for wound healing.

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#### DESCRIPTION

#### TRUNCATED VEGF-RELATED PROTEINS

#### Field Of The Invention

The present invention relates to novel truncated forms of vascular endothelial growth factor (VEGF)-related proteins. More particularly, the invention relates to N-terminally truncated VEGF-related proteins that are substantially free of other proteins. Such truncated VEGF-related proteins may be used to stimulate angiogenesis in vivo and in vitro.

The invention also relates to nucleic acids encoding such novel truncated VEGF-related proteins, cells, tissues and animals containing such nucleic acids; methods of treatment using such nucleic acids; and methods relating to all of the foregoing.

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#### Background

Vascular endothelial growth factors (VEGFs), also called vascular permeability factors (VPFs), are a family of proteins that are produced by many different cell types in many organs 20 and act in a highly selective manner to stimulate endothelial cells almost exclusively (reviewed in Ferrara et al., Endocr. Rev. 13:18-32, (1992); Dvorak et al., Am. J. Pathol. 146:1029-39, 1995; Thomas, J. Biol. Chem. 271:603-06, 1996). These publications, and all other publications referenced herein, are hereby incorporated by reference in their entirety.

When tested in cell culture, VEGFs are potently mitogenic (Gospodarowicz et al., Proc. Natl. Acad. Sci. USA 86:7311-15, 1989) and chemotactic (Favard et al., Biol. Cell 73:1-6, 1991). Additionally, VEGFs induce plasminogen activator, plasminogen activator inhibitor, and plasminogen activator receptor

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also stimulate the formation of tube-like structures by endothelial cells, an in vitro example of angiogenesis (Nicosia et al., Am. J. Pathol., 145:1023-29, 1994).

In vivo, VEGFs induce angiogenesis (Leung et al., Science 5 246:1306-09, 1989) and increase vascular permeability (Senger et al., Science 219:983-85, 1983). VEGFs are now known as important physiological regulators of capillary blood vessel formation. They are involved in the normal formation of new capillaries during organ growth, including fetal growth (Peters et al., Proc. Natl. Acad. Sci. USA 90:8915-19, 1993), tissue 10 repair (Brown et al., <u>J. Exp. Med.</u> 176:1375-79, 1992), the menstrual cycle, and pregnancy (Jackson et al., Placenta 15:341-53, 1994; Cullinan & Koos, <u>Endocrinology</u> 133:829-37, 1993; Kamat et al., Am. J. Pathol. 146:157-65, 1995). During fetal development, VEGFs appear to play an essential role in the de novo formation of blood vessels from blood islands (Risau & Flamme, Ann. Rev. Cell. Dev. Biol. 11:73-92, 1995), as evidenced by abnormal blood vessel development and lethality in embryos lacking a single VEGF allele (Carmeliet et al., Nature 20 380:435-38, 1996). Moreover, VEGFs are strongly implicated in the pathological blood vessel growth characteristic of many diseases, including solid tumors (Potgens et al., Biol. Chem. Hoppe-Seyler 376:57-70, 1995), retinopathies (Miller et al., Am. J. Pathol. 145:574-84, 1994; Aiello et al., N. Engl. J. Med. 331:1480-87, 1994; Adamis et al., Am. J. Ophthalmol. 25 118:445-50, 1994), psoriasis (Fetmar et al., J. Exp. Med. 180:1141-46, 1994), and rheumatoid arthritis (Fava et al., J. Exp. Med. 180:341-46, 1994).

VEGF expression is regulated by hormones (Schweiki et al., 30 <u>J. Clin. Invest.</u> 91:2235-43, 1993) growth factors (Thomas, <u>J. Biol. Chem. 271:603-06, 1996</u>), and by hypoxia (Schweiki et al.,

tissues increase oxygenation through induction of additional capillary vessel formation and resulting increased blood flow. This mechanism is thought to contribute to pathological angiogenesis in tumors and in retinopathies. However, upregulation of VEGF expression after hypoxia is also essential in tissue repair, e.g., in dermal wound healing (Frank et al., J. Biol. Chem. 270:12607-613, 1995), and in coronary ischemia (Banai et al., Cardiovasc. Res. 28:1176-79, 1994; Hashimoto et al., Am. J. Physiol. 267:H1948-H1954, 1994).

10 The potential of VEGF to pharmacologically induce angiogenesis in animal models of vascular ischemia has been shown in the rabbit chronic limb ischemia model demonstrating that repeated intramuscular injection or a single intra-arterial bolus of VEGF can augment collateral blood vessel formation as evidenced by blood flow measurement in the 15 ischemic hindlimb (Pu, et al., Circulation 88:208-15, 1993; Bauters et al., Am. J. Physiol. 267:H1263-71, 1994; Takeshita et al., Circulation 90 [part 2], II-228-34, 1994; Bauters et al., J. Vasc. Surg. 21:314-25, 1995; Bauters et Circulation 91:2802- 09, 1995; Takeshita et al., <u>J. Clin.</u> 20 Invest. 93:662-70, 1994). In this model, VEGF has also been shown to act synergistically with basic FGF to ameliorate ischemia (Asahara et al., Circulation 92: [suppl 2], II-365-71, 1995). VEGF was also reported to accelerate the repair of 25 balloon-injured rat carotid artery endothelium thereby inhibiting pathological thickening of the underlying smooth muscle layers, and thus maintaining lumen diameter and blood flow (Asahara et al., Circulation 91:2793-2801, 1995). has also been shown to induce EDRF (Endothelium-Derived 30 Relaxing Factor (nitric oxide))-dependent relaxation in canine

coronary arteries, thus potentially contributing to increased

for a potential therapeutic role of VEGFs in wound healing, ischemic diseases and restenosis.

The VEGF family of proteins is comprised of at least 4 members VEGF-121, VEGF-165, VEGF-189, and VEGF-206. originally characterized VEGF is a 34-45 kDa glycosylated protein which consists of 2 identical subunits of 165 amino acid residues (Tischer et al., Biochem. Biophys. Res. Commun. 165:1198-1206, 1989). The VEGF-165 cDNA encodes a 191-residue amino acid sequence consisting of a 26-residue secretory signal peptide sequence, which is cleaved upon secretion of the 10 protein from cells, and the 165-residue mature protein subunit. VEGF-165 binds strongly to hoparin for which the strongly basic sequence between residues 115-159 is thought to be responsible (Fig. 1) (Thomas, J. Biol. Chem., 271:603-06 (1996)). other members of the VEGF family are homodimeric proteins with shorter or longer subunits of 121, 189 and 206 residues (VEGF-121, VEGF-189, and VEGF-206, respectively) (Tischer et al., J. Biol. Chem. 266:11947-54, 1991; Park et al., Mol Biol Cell 4:1317-26 (1993)). The 4 forms of VEGF arise from alternative 20 splicing of up to 8 exons of the VEGF gene (VEGF-121, exons 1-5,8; VEGF-165, exons 1-5,7,8; VEGF-189, exons 1-5, 6a, 7, 8; VEGF-206, exons 1-5, 6b, 7, 8 (exon 6a and 6b refer to 2 alternatively spliced forms of the same exon)) (Houck et al., Mol. Endocr., 5:1806-14 (1991)). The VEGF sequences contain 25 eight conserved disulfide-forming core cysteine residues. All VEGF genes encode signal peptides that direct the protein into the secretory pathway. However, only VEGF-121 and -165 are found to be readily secreted by cultured cells whereas VEGF-189 and -206 remain associated with the extracellular matrix. These VEGF forms possess an additional highly basic sequence,

30 corresponding to residues 115-139 in VEGF-189 and -206 (matrix-

Mitogenic activity of the various VEGF isoforms varies depending on each isoform. For example, VEGF-121 and VEGF-165 have very similar mitogenic activity for endothelial cells. However, VEGF-189 and VEGF-206 are only weakly mitogenic (Ferrara et al., Endocr. Rev. 13:18-32, 1992). The reduced activity of these isoforms is attributed to their strong association with cells and matrix, as evidenced by the normal mitogenic activity of a mutant of VEGF-206 which lacks the 24-residue "matrix targeting" sequence common to VEGF-189 and VEGF-206 (residues 115-139 in Fig. 1) (Ferrara et al., Endocr. Rev. 13:18-32, 1992).

An N-terminal fragment of VEGF-165 generated by plasmin (VEGF (1-110)) bound with the same affinity to the KDR receptor as VEGF-165 and VEGF-121 whereas the C-terminal VEGF-fragment (111-165) had no binding activity (Keyt et al., J. Biol. Chem. 15 271:7788-95, 1996). Interestingly, in this study the mitogenic activity of VEGF-121 and VEGF-110 was reduced by approximately 110-fold as compared to VEGF-165, suggesting a potential role of the C-terminal domain of VEGF-165 in the biological potency of VEGF isoforms. The significance of this finding is somewhat 20 unclear in view of earlier results showing the equivalent potency of VEGF-121 and VEGF-165 on endothelial cell growth. Furthermore, since functional interaction of VEGF with the KDR receptor is thought to be dependent at least in part on cell 25 surface heparin sulfate proteoglycan(s) (Cohen et al., J. Biol. Chem., 270:11322-26, 1995; Tessler et al., J. Biol. Chem. 269:12456-61; 1994) it is conceivable that differences in results arise from differences in various experimental systems. In this context it is unclear to what extent cell surface 30 heparin sulfates regulate the functional interaction of VEGF-121 (lacking a heparin-binding domain) and VEGF-165 (possessing

VEGFs are related to platelet-derived growth factor (PDGF) (Andersson et al., Growth Factors 12:159-64, 1995). VEGFs are also related to the family of proteins derived from the Placenta Growth Factor (PIGF) gene, PIGF-129 and PIGF-150 (Maglione et al., Proc. Natl. Acad. Sci. USA 88:9267-71, 1991; Oncogene 8:925-31, 1993). More recently several additional VEGF-related genes have been identified and termed VEGF-B (also called VEGF-related factor VRF-1) (Grimmond et al., Genome Res. 6:122-29, 1996; Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 10 93:2567-81, 1996) VRF-2 (Grimmond et al., Genome Res. 6:122-29, 1996), and VEGF-C (Joukov et al., EMBO J. 15:290-98, 1996; Lee et al., Proc. Natl. Acad. Sci. USA 93:1968-92, 1996) and VEGF-3 (PCT Application No. PCT/US95/07283, published on December 12, 1996 as WO96/39421). Finally, two virally encoded VEGF-related sequences have been identified, poxvirus ORF-1 and ORF-2 15 (Lyttle et al., J. Virol. 68:84-92, 1994). With the exception of PDGF, these proteins are referred to as VEGF-related proteins [VRPs]. Sequences of examples of VRPs are depicted in Figure 1.

The VRPs, and the PDGFs known so far have 8 cysteines within their sequences that are relatively positionally conserved. The protein sequence spanning the conserved cysteines is therefore referred to herein as the core sequence, and the first N-terminal conserved cysteine of the sequence is referred to herein as the "First cysteine of the core sequence" or "first core cysteine."

Interestingly, members of the VEGF families can form heterodimers, such as heterodimers consisting of VEGF and PlGF subunits (DiSalvo et al., <u>J. Biol. Chem.</u> 270:7717-23, 1995; Cao et al., <u>J. Biol. Chem.</u> 271: 3154-62, 1996). Whereas VEGFs are highly potent in stimulating angiogenesis and endothelial cell

<u>J. Biol. Chem.</u> 271: 3154-62, 1996). In other experiments, VEGF-165/VEGF-B heterodimers were found to form after transfection of cells with both genes (Olofsson et al., <u>Proc.</u> Natl. Acad. Sci. U.S.A. 93:2567-81, 1996).

VEGFs interact with two receptors present on endothelial cells, KDR/flk-1 (Terman et al., <u>Biochem. Biophys. Res. Commun.</u> 187:1579-86, 1992), and flt-1 (De Vries et al., <u>Science 255:989-91, 1992)</u>. Systematic site-directed mutagenesis of VEGF-165 by alanine scanning of charged residues has shown that residues D63, E64 and E67 are involved in binding of VEGF to flt-1 whereas the basic residues R82, KI84, and H86 contribute strongly to binding to KDR (Keyt et al., <u>J. Biol. Chem.</u> 271:5638-46, 1996).

VRPs are known to bind to one or more of three different endothelial cell receptors, each of which is a single 15 transmembrane protein with a large extracellular portion comprised of 7 immunoglobulin-type domains and a cytoplasmic portion that functions as a tyrosine kinase. These receptors are KDR/flk-1 (Terman et al., Biochem. Biophys. Res. Commun. 20 187:1579-86, 1992), flt-1 (De Vries et al., Science 255:989-91, 1992), and flt-4 (Pajusola et al., Cancer Res. 52:5738-43, There are distinct selectivities between these receptors and the various VEGF ligands that have not been completely elucidated as yet. However, it is known that VEGF 25 binds to KDR and flt1 (Terman et al., Growth Factors 11:187-95, 1994) but not flt4 (Joukov et al., <u>EMBO J.</u> 15:290-98, 1996), PlGF binds to flt 1 but not KDR (Terman et al., Growth Factors 11:187-95, 1994) and flt4 (Joukov et al., EMBO J. 15:290-98, 1996), VEGF-C binds to flt-4 (Joukov et al., EMBO J. 15:290-98, 30 1996) but it is controversial whether it also binds to KDR (Joukov et al., EMBO J. 15:290-98, 1996; Lee et al., Proc.

endothelial cell proliferation (Olofsson et al., <u>Proc. Natl.</u>
<u>Acad. Sci. U.S.A.</u> 93:2567-81, 1996) it may be speculated that
VEGF-B can bind to KDR because KDR is thought to be primarily
responsible for the angiogenic response of endothelial cells to

VEGF-like growth factors (Gitay-Goren et al., <u>J. Biol. Chem.</u>
271:5519-23 (1996)).

Most of the VRPs have been shown to activate the KDR receptor which is thought to make endothelial cells "angiogenesis-competent." Evidence for such activity has been presented for VEGF-B which stimulates endothelial cell 10 proliferation (Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 93:2567-81, 1996), VEGF-C which stimulates endothelial cell migration and proliferation (Joukov et al., EMBO J. 15:290-98, 1996; Lee et al., Proc. Natl. Acad. Sci. USA 93:1988-92, 1996), and both known virally encoded VRPs which were reported to be 15 angiogenic (Lyttle et al., J. Virol. 68:84-92, 1994). A notable exception are PIGF isoform homodimers which have negligible mitogenic activity for endothelial cells. However, PlGF/VEGF heterodimers still retain considerable mitogenic activity (DiSalvo et al., J. Biol. Chem. 270:7717-23, 1995; Cao 20 et al., J. Biol. Chem. 271: 3154-62, 1996).

VEGFs are expressed in many different tissues. Similarly, VRP genes are also expressed in multiple tissues but it is of particular interest that VEGF-B and to a lesser extent VRF-2 are strongly expressed in human heart and skeletal muscle (Grimmond et al., Genome Res. 6:122-29, 1996; Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 93:2567-81, 1996). In fact, VEGF-B is expressed considerably more strongly in mouse heart tissue than VEGF (Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 93:2567-81, 1996). VEGF-C is also strongly expressed in several human tissues, most notably in heart and skeletal

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physiological role in angiogenesis in these tissues. This is thought to be relevant in pathological situations such as coronary ischemia where collateral angiogenesis is required to provide the heart muscle with an adequate capillary blood vessel supply. It has been shown that transient ischemia induced by coronary artery ligation or hypoxia rapidly upregulates VEGF mRNA in the rat or pig heart in vivo and hypoxia induces VEGF mRNA in cardiac myocytes and smooth muscle cells in vitro (Hashimoto et al., Am. J. Physiol 267, H1948-H1954, 1994; Banai, et al., Cardiovac. Res. 28:1176-79, 1994; 10 Circulation 90, 649-52, 1994). The strong expression of VEGF and VRPs in the heart may help to ensure a redundant and competent regulatory system capable of inducing new blood vessel formation when it is needed. Collateral blood vessel formation is also required in peripheral (lower limb) vascular 15 ischemias and in cerebral ischemias (stroke). Finally, new blood vessel formation is required in tissue repair after wounding. In this context, it is worth noting that VEGF is upregulated in epidermal keratinocytes during skin wound healing (Brown et al., J. Exp. Med. 176:1375-79, 1992). Thus, 20 therapy of various ischemic conditions such as cardiac infarction, chronic coronary ischemia, chronic lower limb ischemia, wound healing and stroke with VRPs may be potentially clinically beneficial.

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#### Summary Of The Invention

The present invention is directed to novel truncated forms of VEGF-related proteins (VRPs), preferably human VRPs. The preferred use of the truncated VRPs and nucleic acid molecule compositions of the invention is to use such compositions to aid in the treatment of patients with heart disease, wounds, or

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VRPs and VEGF proteins (core cysteines). VRPs include, but are not limited to, VEGF-B, VEGF-C, VRF-2, ORF-1, ORF-2, and PlGFs.

A first aspect of the invention provides for a truncated VRP having a deletion of at least one of the amino acid residues N-terminal to the first cysteine of the core sequence of said subunit. Such compositions would be substantially free of other proteins. Preferably, the truncations range from truncating minimally the N-terminal residue of the mature protein subunit only(not including the signal sequence) and maximally all N-terminal amino acids of the mature protein up to and including the residue N-terminal to (prior to) the first core cysteine residue. In more preferred aspects, all of the amino acid residues N-terminal to the first cysteine of the core sequence, except the 1 to 5 amino acid residues immediately N-terminal to said first cysteine, are deleted.

Although the amino acid deletions may consist of deletions of non-adjacent amino acid residues in the N-terminal sequence, it is preferred that the deletions be of consecutive amino acid residues. Thus, in one preferred aspect, the invention comprises human VRPs that have deletions of amino acid residue sequences of increasing lengths from the N-terminus of the N-terminal sequence up to the first cysteine of the core sequence of the VRP subunit sequence.

In preferred aspects, the invention provides for truncated versions of the VRPs VEGF-B, VRF-2, VEGF-C, VEGF-3, PlGF, poxvirus ORF-1, and poxvirus ORF-2. In such truncated VRPs, each subunit may independently have a deletion of at least one of the amino acid residues N-terminal to the first cysteine of the core sequence of said subunit, or only one of the subunits may have such a deletion.

In particular embodiments, the truncated VRP subunit

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consist of consecutive amino acid residues derived from the N-terminal sequence. These consecutive N-terminal residues may be derived from any location in the N-terminal sequence, however, a consecutive sequence starting from the N-terminus of the N-terminal sequence is preferred, and a sequence consisting of consecutive amino acid residues immediately N-terminal to the first cysteine of the core sequence of the VRP subunit is most preferred. Examples of such most preferred embodiments are depicted in Figure 2.

In other embodiments, the amino acid residues N-terminal to the first cysteine of the core sequence of the truncated VRPs of the invention are a randomly selected amino acid sequence, in yet other embodiments, these amino acid residues are derived from the N-terminal sequence of the full length VRP sequence, but are not necessarily consecutive amino acids from the full length VRP sequence.

Thus, in one most preferred aspect, the invention provides a truncated VRP subunit wherein the amino acid residues N-terminal to the first cysteine of the core sequence of said subunit are deleted.

In other aspects, the invention provides a truncated VRP subunit wherein the amino acid sequence N-terminal to the core sequence comprises 11 to 20, more preferably 11 to 15, more preferably 6 to 10, and most preferably 2 to 5 amino acid residues.

Preferably, the amino acid sequence N-terminal to the core sequence comprises the consecutive amino acid residues immediately N-terminal to the first cysteine of the core sequence of said VRP subunit. Thus, in these preferred embodiments, the truncated VRP comprises the core sequence, the necessary C-terminal sequence to the core sequence, and further

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consecutive amino acid residues of the amino acid sequence that is immediately N-terminal to the first cysteine of the core sequence of the full length VRP sequence.

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Those skilled in the art will recognize that where a truncated VRP subunit comprises, for example, (X) amino acids N-terminal to the first cysteine of the core sequence, that such a truncated VRP subunit is one where the corresponding full length VRP subunit comprises (X + 1) amino acids Nterminal to the first cysteine of the core sequence.

The truncated VRPs of the invention include truncated VRP 10 homodimers comprising two truncated VRP subunits of the invention, wherein the two truncated VRP subunits have the same sequence, and also include truncated amino acid heterodimers comprising two truncated VRP subunits of the invention wherein the two subunits have different amino acid 15 sequences from each other.

For purposes of the present invention, the term "first N-NN" amino acids where N and NN each represent numbers of amino acids, for example, the first 10-15 amino acids, denotes the first N-NN amino acids (e.g., the first 10-15 amino acids) after the signal peptide sequence of the designated VRP. term N-NN encompasses a deletion of anywhere from N to NN of the first amino acids after the signal sequence. Thus, in more preferred aspects, the truncated VRP subunit comprises a 25 truncated hVEGFB protein subunit wherein the first 10-15 amino acids are deleted; more preferably, the first 15-20 amino acids are deleted; more preferably, the first 20-25 amino acids are deleted; and most preferably, the first 23-24 amino acids are deleted.

30 In other more preferred aspects, the truncated VRP subunit comprises a truncated hVRF2 protein subunit wherein the first

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acids are deleted; and most preferably, the first 23-24 amino acids are deleted.

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In other more preferred aspects, the truncated VRP subunit comprises a truncated hVEGFC protein subunit wherein the first 95-100 amino acids are deleted; more preferably, the first 100-105 amino acids are deleted; more preferably, the first 105-110 amino acids are deleted; and most preferably, the first 108-109 amino acids are deleted.

In other more preferred aspects, the truncated VRP subunit comprises a truncated hPlGF protein subunit wherein the first 16-21 amino acids are deleted; more preferably, the first 21-26 amino acids are deleted; more preferably, the first 26-31 amino acids are deleted; and most preferably, the first 29-30 amino acids are deleted.

In other more preferred aspects, the truncated VRP subunit comprises a truncated hVEGF3 protein subunit wherein the first 10-15 amino acids are deleted; more preferably, the first 15-20 amino acids are deleted; more preferably, the first 20-25 amino acids are deleted; and most preferably, the first 23-24 amino acids are deleted.

In other more preferred aspects, the truncated VRP subunit comprises a truncated pvORF1 protein subunit wherein the first 20-25 amino acids are deleted; more preferably, the first 25-30 amino acids are deleted; more preferably, the first 30-35 amino acids are deleted; and most preferably, the first 33-34 amino acids are deleted.

In other more preferred aspects, the truncated VRP subunit comprises a truncated pvORF2 protein subunit wherein the first 30-35 amino acids are deleted; more preferably, the first 35-40 amino acids are deleted; more preferably, the first 40-45 amino acids are deleted; and most preferably, the first 43-44 amino

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The invention also provides for nucleic acid molecules coding for the truncated VRP subunits described herein. The nucleic acid molecules may be, for example, DNA, cDNA or RNA. The invention also provides for recombinant DNA vectors comprising the nucleic acid molecules encoding the truncated VRPs, and host cells transformed with such recombinant DNA vectors, wherein such vectors direct the synthesis of a truncated VRP subunit such as those described herein.

The invention further provides for nucleic acid molecules encoding biosynthetic precursor forms of N-terminally truncated subunits of VRPs for the purpose of facilitating the expression in suitable host systems. Such nucleic acid molecules are comprised of DNA encoding a signal peptide that precedes the truncated subunits at their N-termini. The signal sequences of VEGF or VRPs would be used to construct appropriate signal peptide-containing truncated forms of VRPs. The human VEGF signal peptide is as follows:

mnfllswvhwslalllylhhakwsqa (I) -- [SEQ I.D. NO. 40] -- Alternatively, the signal peptides shown in Figure 1 may be used. Preferably, the signal peptide specific for the truncated VRP is used in the construct.

In order to facilitate signal peptide cleavage in mammalian cells after fusion of the signal sequence to truncated forms of VRP, it may be necessary to include the first or the first two residues of the mature VRP peptide sequence, e.g. proline (P), or proline-valine (PV) for hVEGFB. Thus, an appropriate nucleic acid molecule would be comprised of DNA encoding the signal sequence of VEGF-B, optionally followed by a codon for proline (the first residue of mature VEGF-B), optionally followed by a codon for valine (the second residue of mature VEGF-B), and followed by DNA encoding the N-

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the art. Those skilled in the art will recognize that the signal peptides should optionally include residues needed for facilitation of signal peptide cleavage in mammalian cells for the various truncated VRP subunits of the present invention.

Thus, the present invention provides for recombinant DNA 5 expression vectors wherein the 5' end of the nucleic acid molecule coding for the truncated VRP subunit is operably linked to a DNA sequence that codes for a signal peptide. signal peptide may be a human VRP signal peptide. Moreover, the DNA sequence coding for said signal peptide may be operably 10 linked at the 3' end of said DNA sequence to DNA coding for the first amino acid residue of the mature non-truncated VRP subunit, and wherein the 3' end of said DNA coding for said residue is operably linked to the nucleic acid molecule coding for said truncated VRP subunit. In other aspects, the DNA 15 sequence coding for said signal peptide is operably linked at the 3' end of said DNA sequence to DNA coding for the first two amino acid residues of the mature non-truncated VRP subunits, and wherein the 3' end of said DNA coding for said two residues is operably linked to said nucleic acid molecule coding for 20 said truncated VRP subunit. Thus, in preferred aspects, the invention also provides a truncated VRP subunit of invention as described above, further comprising at the Nterminus of said truncated VRP subunit, the first one or two amino acid residues of the mature non-truncated VRP subunit. 25 Those skilled in the art will recognize that such truncated VRP subunits of the invention include those wherein the final number of amino acids N-terminal to the first cysteine of the core sequence (including the additional one or two amino acids that may facilitate signal peptide cleavage) is at least one 30 less than the number of amino acids N-terminal to the first

In other preferred aspects, the invention provides truncated VRP homodimers or heterodimers comprising two truncated VRP subunits wherein said truncated VRP subunits comprise at the N-terminus of said truncated VRP subunits, the first one or two amino acid residues of the mature non-truncated VRP subunit.

In preferred aspects, the recombinant nucleic acid molecule coding for a truncated VRP subunit of the invention is operably linked to control sequences operable in a host cell transformed with said vector. The present invention also provides for transformed or transfected host cells comprising the recombinant DNA vectors of the invention.

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The present invention also includes delivery vectors which comprise nucleic acid molecules coding for the truncated VRPs of the invention. Such delivery vectors may be, for example, viral vectors. Such viral vectors may be, for example, adenovirus vectors or adenovirus-associated virus vectors. other aspects of the invention are provided an adenovirus vector comprising a nucleic acid molecule coding for a truncated VRP of the invention operably linked at the 5' end of the nucleic acid molecule to a DNA sequence that codes for a signal peptide. Preferably, the signal peptide is selected from the group consisting of VEGF signal peptide, VEGF-B signal peptide, VRF-2 signal peptide, VEGF-C signal peptide, PlGF signal peptide, VEGF-3 signal peptide, poxvirus ORF-1 signal peptide, and poxvirus ORF-2 signal peptide. Preferably said signal peptide is VEGF-B signal peptide. In preferred aspects, the DNA sequence coding for the signal peptide is operably linked at the 3' end of the DNA sequence to DNA coding for the first amino acid residue of the mature non-truncated VRP subunit, and wherein the 3' end of said DNA coding for said

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adenovirus vector comprises a nucleic acid molecule which codes for a truncated VRP subunit of Figure 2.

In further preferred aspects of the invention are provided a filtered-injectable adenovirus vector preparation comprising a recombinant adenoviral vector, said vector containing no wild-type virus and comprising: a partial adenoviral sequence from which the E1A/E1B genes have been deleted, and a transgene coding for a truncated VRP subunit, driven by a promoter adenovirus partial sequence; and flanked by the pharmaceutically acceptable carrier. In preferred aspects, the preparation has been filtered through a 30 micron filter. other preferred aspects the truncated VEGF subunit is a truncated VEGF subunit of Figure 2. In another preferred aspect, the injectable adenoviral vector preparation comprises a promoter selected from the group consisting of a  ${\tt CMV}$ promoter, a ventricular myocyte-specific promoter, and a myosin heavy chain promoter.

In other aspects, the invention provides a method of producing a truncated VRP polypeptide comprising growing, under suitable conditions, a host cell transformed or transfected with the recombinant DNA expression vector of the invention in a manner allowing expression of said polypeptide, and isolating said polypeptide from the host cell. Suitable conditions are then provided for the truncated VRP peptide to fold into a truncated VRP subunit. In mammalian cells, such conditions should be naturally provided by the cell. In non-mammalian cells, appropriate pH, isotonicity, and reducing conditions must be provided, such as those described in, for example, Example 2. Most preferably, the invention provides a method of producing a truncated VRP wherein suitable conditions are provided for said truncated VRP subunit to dimerize with a

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homodimer comprising two truncated VRP subunits having the same amino acid sequence.

In other aspects of the invention are provided methods of producing truncated VRP heterodimers wherein the two VRP amino have different acid sequences. Such subunits heterodimers may consist of one truncated VRP subunit and one non-truncated VRP subunit, or both VRP subunits The two subunits may be derived from different truncated. VRPs. For example, the heterodimer may consist of one VEGF-B subunit and one truncated VEGF-C subunit, or both subunits may be truncated.

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In further preferred aspects, the present invention provides pharmaceutical compositions comprising a truncated VRP subunit of the present invention, in a suitable carrier. The invention includes methods of stimulating blood vessel formation comprising administering to a patient such a pharmaceutical composition.

Methods are provided using the compounds of the present invention to stimulate endothelial cell growth or endothelial cell migration in vitro comprising treating said endothelial cells with truncated VRPs.

The present invention also provides methods of treating a patient suffering from a heart disease comprising administering to said patient a nucleic acid molecule coding for at least one truncated VRP subunit, said nucleic acid molecule capable of expressing the truncated VRP subunit in said patient. In an additional embodiment, methods are provided of stimulating angiogenesis in a patient comprising administering a therapeutic amount of a pharmaceutical composition comprising a truncated VRP of the present invention.

Preferably, the pharmaceutical composition is in a

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for example, basic Fibroblast Growth Factor (bFGF) (FGF-2), acidic FGF (aFGF) (FGF-1), FGF-4, FGF-5, FGF-6, or any FGF or other angiogenic factor that stimulates endothelial cells. one aspect of the invention is provided Thus, in pharmaceutical composition comprising a truncated VRP and one or more potentiating agents. The pharmaceutical compositions may also be used to treat patients suffering from ischemic conditions such as cardiac infarction, chronic coronary ischemia, chronic lower limb ischemia, stroke, and peripheral Methods are also provided using the vascular disease. 10 pharmaceutical compositions of the present invention to treat wounds, such as dermal or intestinal wounds.

In preferred embodiments, methods are provided of stimulating angiogenesis in a patient comprising delivering a delivery vector to the myocardium of the patient by intracoronary injection directly into one or both coronary arteries, said vector comprising a nucleic acid molecule coding for at least one truncated VRP subunit, wherein said vector is capable of expressing the truncated VRP subunit in the myocardium.

In other preferred embodiments, the method may be used for stimulating coronary collateral vessel development.

In more preferred embodiments, a method is provided for stimulating vessel development in a patient having peripheral vascular disease, comprising delivering a delivery vector to the peripheral vascular system of the patient by intra-femoral artery injection directly into one or both femoral arteries, said vector comprising a transgene coding for a truncated VRP subunit, and capable of expressing the truncated VRP subunit in the peripheral vascular system, thereby promoting peripheral vascular development.

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another preferred aspect, the delivery vector is an adenoassociated virus vector.

#### Brief Description Of The Drawings

Figure 1 depicts the amino acid sequences of VEGF-B [SEQ I.D. NO. 1], VRF-2 [SEQ I.D. NO. 2], VEGF-C [SEQ I.D. NO. 3], PlGF (human PlGF-2) [SEQ I.D. NO. 4], VEGF-3 [SEQ I.D. NO. 5], poxvirus ORF-1 [SEQ I.D. NO. 6], and poxvirus ORF-2 [SEQ I.D. NO. 7]. Lower case letters signify signal peptides that are cleaved from the mature protein. The eight cysteines of the 10 core sequence are underlined. Sequences are described in the following references: human VEGF-B: Grimmond et al., Genome Res. 6:122-29 (1996); Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 93:2567-81 (1996); mouse VEGF-B: Olofsson et al., Proc. Natl. Acad. Sci. U.S.A. 93:2567-81 (1996); human VRF-2: 15 Grimmond et al., Genome Res. 6:122-29 (1996); human VEGF-C: Joukov et al., EMBO J. 15:290-98 (1996); Lee et al., Proc. Natl. Acad. Sci. USA 93:1988-92 (1996); PlGF: Maglione et al., Oncogene 8:925-31 (1993); Hauser & Weich, Growth Factors 9:259-68 (1993); human VEGF3: PCT Application Serial No. 20 PCT/US95/07283, published on December 12,, 1996, as WO96/39421; poxvirus ORF-1 and ORF-2: Lyttle et al., J. Virol. 68:84-92 (1994).

Figure 2a-2f depicts examples of truncated VRP amino acid sequences below the corresponding full length (F/L) VRP sequence. The amino acid sequences of each truncation are listed as follows:

2a(F/L)[SEQ I.D. NO. 34](1) [SEQ I.D. NO. 8]; 2a(2) [SEQ I.D. NO. 9]; 2a(3) [SEQ I.D. NO. 10]; 2a(4) [SEQ I.D. NO. 11]; 30 2a(5) [SEQ I.D. NO. 12]; 2a(6) [SEQ I.D. NO. 13]; 2b (F/L) [SEQ I.D. NO. 35]; (1) [SEQ I.D. NO. 14]; 2b(2) [SEQ I.D. NO. 15];

2c(2) [SEQ I.D. NO. 19]; 2c(3) [SEQ I.D. NO. 20]; 2c(4) [SEQ I.D. NO. 21]; 2d(F/L) [SEQ I.D. NO. 37]; (1) [SEQ I.D. NO. 22]; 2d(2) [SEQ I.D. NO. 23]; 2d(3) [SEQ I.D. NO. 24]; 2d(4) [SEQ I.D. NO. 25]; 2e(F/L [SEQ I.D. NO. 38] (1) [SEQ I.D. NO. 26]; 2e(2) [SEQ I.D. NO. 27]; 2e(3) [SEQ I.D. NO. 28]; 2e(4) [SEQ I.D. NO. 29]; 2f(F/L) [SEQ I.D. NO. 39]; (1) [SEQ I.D. NO. 30]; 2f(2) [SEQ I.D. NO. 31]; 2f(3) [SEQ I.D. NO. 32]; and 2f(4) [SEQ I.D. NO. 33].

# 10 <u>Detailed Description Of The Invention</u> Construction of Novel Truncated VRP Sequences

In a first aspect the invention features a truncated VRP comprising at least one truncated VRP subunit. By "truncated VRP subunit" it is meant a VRP subunit having an amino acid sequence substantially similar to one of the VRPs, for example, 15 but not limited to, one of the sequences shown in Figure 1, or an analog or derivative thereof, wherein at least one of the Nterminal amino-acid residues N-terminal to the first cysteine of the core sequence of the mature subunit is deleted. A sequence that is "substantially similar" to a VRP comprises an 20 amino acid sequence that is at least 25% homologous to the 8 cysteine core sequence of VEGF-B, comprises all of the essential conserved cysteine residues of said core sequence, and retains VRP activity. By "truncated VRP subunit" is also meant a VRP subunit wherein at least one of the N-terminal 25 amino acid residues N-terminal to the first cysteine of the VEGF core sequence is deleted, and, at least one of the cysteines of the core sequence is deleted, wherein said cysteine is non-essential. A non-essential cysteine is one that is not required to retain VRP activity. Such non-30 essential cysteines have been described in connection with

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measured by dividing the number of identical residues by the total number of residues and multiplying the product by 100. Thus, two copies of exactly the same sequence have 100% identity, but sequences that are less nightly conserved and have deletions, additions, or replacements may have a lower degree of identity. In calculating sequence identity, the two sequences are compared starting at the carboxy terminus of the N-terminal deletion. Those skilled in the art will recognize that several computer programs are available for determining sequence identity.

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Analogs of a truncated VRP polypeptide or subunit are functional equivalents having similar amino acid sequence and retaining, to some extent, one or more activities of the related truncated VRP polypeptide or subunit. By "functional equivalent" is meant the analog has an activity that can be substituted for one or more activities of a particular truncated VRP polypeptide or subunit. Preferred functional equivalents retain all of the activities of a particular truncated VRP polypeptide or subunit, however, the functional an activity that, when measured may have equivalent quantitatively, is stronger or weaker, as measured in VRP functional assays, for example, such as those disclosed herein. In most cases, such truncated VRP polypeptides or subunits must be incorporated into a truncated VRP dimer in order to measure functional activity. Preferred functional equivalents have activities that are within 1% to 10,000% of the activity of the related truncated VRP polypeptide or subunit, more preferably between 10% to 1000%, and more preferably within 50% to 200%.

The ability of a derivative to retain some activity can be 30 measured using techniques described herein. Derivatives include modification occurring during or after translation, for

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molecule, membrane molecule or other ligand (see Ferguson et al., 1988, Annu. Rev. Biochem. 57:285-320).

Specific types of derivatives or analogs also include amino acid alterations such as deletions, substitutions, additions, and amino acid modifications. A "deletion" refers to the absence of one or more amino acid residue(s) in the related polypeptide. An "addition" refers to the presence of one or more amino acid residue(s) in the related polypeptide. Additions and deletions to a polypeptide may be at the amino terminus, the carboxy terminus, and/or internal. Amino acid "modification" refers to the alteration of a naturally occurring amino acid to produce a non-naturally occurring amino acid. A "substitution" refers to the replacement of one or more amino acid residue(s) by another amino acid residue(s) in Derivatives can contain different the polypeptide. combinations of alterations including more than one alteration and different types of alterations.

While the effect of an amino acid change on VRP activity varies depending upon factors such as phosphorylation, glycosylation, intra-chain linkages, tertiary structure, and the role of the amino acid in the active site or a possible allosteric site, it is generally preferred that the substituted amino acid is from the same group as the amino acid being replaced. To some extent the following groups contain amino acids which are interchangeable: the basic amino acids lysine, arginine, and histidine; the acidic amino acids aspartic and glutamic acids; the neutral polar amino acids serine, threonine, cysteine, glutamine, asparagine and, to a lesser extent, methionine; the nonpolar aliphatic amino acids glycine, alanine, valine, isoleucine, and leucine (however, because of size, glycine and alanine are more closely related and valine,

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alanine, glycine, and serine seem to be interchangeable to some extent, and cysteine additionally fits into this group, or may be classified with the polar neutral amino acids.

Preferred derivatives have one or more amino acid alteration(s) which do not significantly affect the activity of the related truncated VRP polypeptide or subunit. In regions of the truncated VRP polypeptide or subunit not necessary for VRP activity, amino acids may be deleted, added or substituted with less risk of affecting activity. In regions required for VRP activity, amino acid alterations are less preferred as there is a greater risk of affecting VRP activity. Such alterations should be conservative alterations. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent.

Conserved regions tend to be more important for protein activity than non-conserved regions. Standard procedures can be used to determine the conserved and non-conserved regions important for VRP activity using *in vitro* mutagenesis techniques or deletion analyses and measuring VRP activity as described by the present disclosure.

Derivatives can be produced using standard chemical techniques and recombinant nucleic acid molecule techniques. Modifications to a specific polypeptide may be deliberate, as through site-directed mutagenesis and amino acid substitution during solid-phase synthesis, or may be accidental such as through mutations in hosts which produce the polypeptide. Polypeptides including derivatives can be obtained using standard techniques such as those described in Sambrook et al., Molecular Cloning, Cold Spring Harbor Laboratory Press (1989). For example, Chapter 15 of Sambrook describes procedures for

of the invention, or a functional analog or derivative thereof as described herein. The term "truncated VRP polypeptide" also includes a truncated VRP subunit; the term subunit generally referring to a peptide that has been folded into an active three-dimensional structure.

By "truncated VRP" is meant a dimer of two VRP subunits. The two subunits may be derived from two different VRPs where both subunits are truncated VRP subunits. One or both of the subunits may be truncated; the two subunits may also have different N-terminal deletions.

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It is advantageous that the truncated VRP, truncated VRP subunit, or truncated VRP polypeptide be enriched or purified. By the use of the term "enriched" in this context is meant that the specific amino acid sequence constitutes a significantly higher fraction (2 - 5 fold) of the total of amino acid sequences present in the cells or solution of interest than in normal or diseased cells or in the cells from which the sequence was taken. This could be caused by a person by preferential reduction in the amount of other amino acid sequences present, or by a preferential increase in the amount of the specific amino acid sequence of interest, or by a combination of the two. However, it should be noted that enriched does not imply that there are no other amino acid sequences present, just that the relative amount of the sequence of interest has been significantly increased. term "significant" here is used to indicate that the level of increase is useful to the person making such an increase, and generally means an increase relative to other amino acid sequences of about at least 2 fold, more preferably at least 5 to 10 fold or even more. The term also does not imply that there is no amino acid sequence from other sources. The other

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situations in which man has intervened to elevate the proportion of the desired amino acid sequence.

It is also advantageous for some purposes that an amino acid sequence be in purified form. The term "purified" in reference to a polypeptide does not require absolute purity (such as a homogeneous preparation); instead, it represents an indication that the sequence is relatively purer than in the natural environment (compared to the natural level this level should be at least 10 fold greater, e.g., in terms of mg/ml). Purification of at least one order of magnitude, preferably two or three orders, and more preferably four or five orders of magnitude is expressly contemplated. The substance is preferably free of contamination at a functionally significant level, for example 90%, 95%, or 99% pure.

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In another aspect the invention features a nucleic acid molecule encoding a truncated VRP polypeptide or subunit.

In some situations it is desirable for such nucleic acid molecule to be enriched or purified. By the use of the term "enriched" in reference to nucleic acid molecule is meant that the specific DNA or RNA sequence constitutes a significantly higher fraction (2 - 5 fold) of the total DNA or RNA present in the cells or solution of interest than in normal or diseased cells or in the cells from which the sequence was taken. This could be caused by a person by preferential reduction in the amount of other DNA or RNA present, or by a preferential increase in the amount of the specific DNA or RNA sequence, or by a combination of the two. However, it should be noted that enriched does not imply that there are no other DNA or RNA sequences present, just that the relative amount of the sequence of interest has been significantly increased. The term significant here is used to indicate that the level of

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even more. The term also does not imply that there is no DNA or RNA from other sources. The other source DNA may, for example, comprise DNA from a yeast or bacterial genome, or a cloning vector such as pUC19. This term distinguishes from naturally occurring events, such as viral infection, or tumor type growths, in which the level of one mRNA may be naturally increased relative to other species of mRNA. That is, the term is meant to cover only those situations in which a person has intervened to elevate the proportion of the desired nucleic acid.

It is also advantageous for some purposes that a nucleotide sequence be in purified form. The term "purified" in reference to nucleic acid molecule does not require absolute purity (such as a homogeneous preparation); instead, it represents an indication that the sequence is relatively purer than in the natural environment (compared to the natural level this level should be at least 2-5 fold greater, e.g., in terms of mg/ml).

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existing VRP nucleotide sequence by modification using, for example, oligonucleotide site-directed mutagenesis, or by deleting sequences using restriction enzymes, or as described herein. Standard recombinant techniques for mutagenesis such as in vitro site-directed mutagenesis (Hutchinson et al., J. Biol. Chem. 253:6551, (1978), Sambrook et al., Chapter 15, supra), use of TAB® linkers (Pharmacia), and PCR-directed mutagenesis can be used to create such mutations. The nucleic acid molecule may also be synthesized by the triester method or by using an automated DNA synthesizer.

The invention also features recombinant DNA vectors and recombinant DNA expression vectors preferably in a cell or an

transcription in a host cell. The recombinant DNA vector can contain a transcriptional initiation region functional in a cell and a transcriptional termination region functional in a cell.

The present invention also relates to a cell or organism that contains the above-described nucleic acid molecule or recombinant DNA vector and thereby is capable of expressing a truncated VRP peptide. The polypeptide may be purified from cells which have been altered to express the polypeptide. A cell is said to be "altered to express a desired polypeptide" when the cell, through genetic manipulation, is made to produce a protein which it normally does not produce or which the cell normally produces at lower levels. One skilled in the art can readily adapt procedures for introducing and expressing either genomic, cDNA, or synthetic sequences into either eukaryotic or prokaryotic cells.

A nucleic acid molecule, such as DNA, is said to be "capable of expressing" a polypeptide if it contains nucleotide sequences which contain transcriptional and translational regulatory information and such sequences are "operably linked" to nucleotide sequences which encode the polypeptide. precise nature of the regulatory regions needed for gene sequence expression may vary from organism to organism, but general include a promoter region shall in prokaryotes, contains both the promoter (which directs the initiation of RNA transcription) as well as the DNA sequences which, when transcribed into RNA, will signal synthesis initiation. Such regions will normally include those 5'-noncoding sequences involved with initiation of transcription and translation, such as the TATA box, capping sequence, CAAT sequence, and the like.

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for such expression: (1) an exogenous promoter sequence (2) a ribosome binding site (3) a polyadenylation signal (4) a secretion signal. Modifications can be made in the 5'-untranslated and 3'-untranslated sequences to improve expression in a prokaryotic or eukaryotic cell; or codons may be modified such that while they encode an identical amino acid, that codon may be a preferred codon in the chosen expression system. The use of such preferred codons is described in, for example, Grantham et al., Nuc. Acids Res., 9:43-74 (1981), and Lathe, J. Mol. Biol., 183:1-12 (1985) hereby incorporated by reference herein in their entirety.

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If desired, the non-coding region 3' to the genomic VRP sequence may be operably linked to the nucleic acid molecule encoding such VRP subunit. This region may be used in the recombinant DNA vector for its transcriptional termination regulatory sequences, such as termination and polyadenylation. Thus, by retaining the 3'-region naturally contiguous to the DNA sequence encoding a VRP gene, the transcriptional termination signals may be provided. Alternatively, a 3' region functional in the host cell may be substituted.

An operable linkage is a linkage in which the regulatory DNA sequences and the DNA sequence sought to be expressed are connected in such a way as to permit gene sequence expression. Two DNA sequences (such as a promoter region sequence and a truncated VRP sequence) are said to be operably linked if the nature of the linkage between the two DNA sequences does not (1) result in the introduction of a frame-shift mutation in the coding sequence, (2) interfere with the ability of the promoter region sequence to direct the transcription of a truncated VRP gene sequence, or (3) interfere with the ability of the a truncated VRP gene sequence to be transcribed by the promoter

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a truncated VRP gene, transcriptional and translational signals recognized by an appropriate host are necessary.

## Expression and Purification of Novel Truncated VRP Sequences

Examples 2 and 3 describe the expression and purification of novel truncated VRP sequences of the present invention as expressed in baculovirus systems. Those skilled in the art will recognize that the truncated VRPs of the present invention may also be expressed in other cell systems, both prokaryotic and eukaryotic, all of which are within the scope of the present invention. Examples 4-6 provide examples of suitable assays for functional activity of the novel truncated VRPs.

Although the truncated VRPs of the present invention may be expressed in prokaryotic cells, which are generally very efficient and convenient for the production of recombinant proteins, the truncated VRPs produced by such cells will not be glycosylated and therefore may have a shorter half-life in vivo. Prokaryotes most frequently are represented by various strains of E. coli. However, other microbial strains may also be used, including other bacterial strains. Recognized prokaryotic hosts include bacteria such as E. coli, Bacillus, Streptomyces, Pseudomonas, Salmonella, Serratia, and the like. The prokaryotic host must be compatible with the replicon and control sequences in the expression plasmid.

In prokaryotic systems, plasmid vectors that contain replication sites and control sequences derived from a species compatible with the host may be used. Examples of suitable plasmid vectors may include pBR322, pUC118, pUC119 and the like; suitable phage or bacteriophage vectors may include γgt10, γgt11 and the like; and suitable virus vectors may include pMAM-neo, pKRC and the like. Preferably, the selected vector of the present invention has the capacity to replicate in the selected

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necessary to operably link the truncated VRP sequence to a functional prokaryotic promoter. Such promoters may be either constitutive or, more preferably, regulatable (i.e., inducible or derepressible). Examples of constitutive promoters include the int promoter of bacteriophage  $\lambda$ , the bla promoter of the  $\beta$ -lactamase gene sequence of pBR322, and the CAT promoter of the chloramphenicol acetyl transferase gene sequence of pPR325, and the like. Examples of inducible prokaryotic promoters include the major right and left promoters of bacteriophage  $\lambda$ ( $P_L$  and  $P_R$ ), the trp, recA,  $\lambda$ acZ,  $\lambda$ acI, and gal promoters of E. coli, the  $\alpha$ -amylase (Ulmanen et al., J. Bacteriol. 162:176-182(1985)) and the  $\varsigma-28$ -specific promoters of B. subtilis (Gilman et at., Gene sequence 32:11-20(1984)), the promoters of the bacteriophages of Bacillus (Gryczan, In: The Molecular Biology of the Bacilli, Academic Press, Inc., NY (1982)), and Streptomyces promoters (Ward et at., Mol. Gen. Genet. 203:468-478(1986)). Prokaryotic promoters are reviewed by Glick (J. Ind. Microbiot. 1:277-282(1987)); Cenatiempo (Biochimie 68:505-516(1986)); and Gottesman (Ann. Rev. Genet. 18:415-442 (1984)).

Proper expression in a prokaryotic cell also requires the presence of a ribosome binding site upstream of the gene sequence-encoding sequence. Such ribosome binding sites are disclosed, for example, by Gold et at. (Ann. Rev. Microbiol. 35:365-404(1981)). The ribosome binding site and other sequences required for translation initiation are operably linked to the nucleic acid molecule coding for the truncated by, for example, in frame ligation of synthetic VRP oligonucleotides that contain such control sequences. expression in prokaryotic cells, no signal peptide sequence is 30 required. The selection of control sequences, expression vectors, transformation methodo and the like and the like

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Thus, the words "transformants" or "transformed progeny. cells" include the primary subject cell and cultures derived therefrom, without regard to the number of transfers. Truncated VRP peptides expressed in prokaryotic cells expected to comprise a mixture of properly truncated VRP peptides with the N-terminal sequence predicted from sequence of the expression vector, and truncated VRP peptides which have an N-terminal methionine resulting from inefficient cleaving of the initiation methionine during bacterial 10 expression. Both types of truncated VRP peptides considered to be within the scope of the present invention as the presence of an N-terminal methionine is not expected to affect biological activity. It is also understood that all progeny may not be precisely identical in DNA content, due to 15 deliberate or inadvertent mutations. However, as defined, mutant progeny have the same functionality as that of the originally transformed cell.

Preferred prokaryotic vectors include plasmids such as those capable of replication in E. coil (such as, for example, pBR322, ColEl, pSC101, pACYC 184,  $\pi$ VX. Such plasmids are, for 20 example, disclosed by Sambrook (cf. "Molecular Cloning: A Laboratory Manual", second edition, edited by Sambrook, Fritsch, & Maniatis, Cold Spring Harbor Laboratory, (1989)). Bacillus plasmids include pC194, pC221, pT127, and the like. 25 Such plasmids are disclosed by Gryczan (In: The Molecular Biology of the Bacilli, Academic Press, NY (1982), pp. 307-329). Suitable Streptomyces plasmids include p1J101 (Kendall et al., J. Bacteriol. 169:4177-4183 (1987)), and streptomyces bacteriophages such as  $\Phi$ C31 (Chater et al., In: 30 International Symposium on Actinomycetales Biology, Akademiai Kaido, Budapest, Hungary (1986), pp. 45-54). Pseudomonas WO 98/49300 PCT/US98/07801

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Eukaryotic host cells which may be used in the expression systems of the present invention are not strictly limited, provided that they are suitable for use in the expression of the truncated VRP peptide. Preferred eukaryotic hosts include, for example, yeast, fungi, insect cells, mammalian cells either in vivo, or in tissue culture. Mammalian cells which may be useful as hosts include HeLa cells, cells of fibroblast origin such as VERO or CHO-K1, or cells of lymphoid origin and their derivatives.

The truncated VRPs of the present invention may also be expressed in human cells such as human embryo kidney 293EBNA cells which express Epstein-Barr virus nuclear antigen 1, as described, for example, in Olofsson, B. et al., Proc. Natl. Acad. Sci. USA 93:2576-2581 (1996). The cells are transfected with the expression vectors of Example 2 by using calcium phosphate precipitation, and the cells are then incubated for at least 48 hours. The truncated VRP peptides may then be purified from the supernatant as described in Example 3.

In addition, plant cells are also available as hosts, and control sequences compatible with plant cells are available, such as the cauliflower mosaic virus 35S and 19S, and nopaline synthase promoter and polyadenylation signal sequences. Another preferred host is an insect cell, for example the Drosophila larvae. Using insect cells as hosts, the Drosophila alcohol dehydrogenase promoter can be used. Rubin, Science 240:1453-1459(1988).

Any of a series of yeast gene sequence expression systems can be utilized which incorporate promoter and termination elements from the actively expressed gene sequences coding for glycolytic enzymes are produced in large quantities when yeast are grown in mediums rich in glucose. Known glycolytic gene

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A number of recombinant DNA strategies exist which utilize strong promoter sequences and high copy number of plasmids which can be utilized for production of the desired proteins in yeast. Yeast recognizes leader sequences on cloned mammalian gene sequence products and secretes peptides bearing leader sequences (i.e., pre-peptides). For a mammalian host, several possible vector systems are available for the expression of truncated VRP peptides.

A wide variety of transcriptional and translational regulatory sequences may be employed, depending upon the nature 10 of the host. The transcriptional and translational regulatory signals may be derived from viral sources, such as adenovirus, bovine papilloma virus, cytomegalovirus, simian virus, or the like, where the regulatory signals are associated with a particular gene sequence which has a high level of expression. Alternatively, promoters from mammalian expression products, such as actin, collagen, myosin, and the like, may be employed. Transcriptional initiation regulatory signals may be selected which allow for repression or activation, so that expression of the gene sequences can be modulated. Of interest are 20 regulatory signals which are temperature-sensitive so that by varying the temperature, expression can be repressed or initiated, or are subject to chemical (such as metabolite) regulation.

Expression of truncated VRPs in eukaryotic hosts requires the use of eukaryotic regulatory regions. Such regions will, in general, include a promoter region sufficient to direct the initiation of RNA synthesis. Preferred eukaryotic promoters include, for example, the promoter of the mouse metallothionein I gene sequence (Hamer et al., <u>J. Mol. Appl. Gen. 1:273-288(1982))</u>; the TK promoter of Herpes virus (McKnight, Cell

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(USA) 79:6971-6975(1982); Silver et al., Proc. Natl. Acad. Sci. (USA) 81:5951-5955 (1984)).

Translation of eukaryotic mRNA is initiated at the codon which encodes the first methionine. For this reason, it is preferable to ensure that the linkage between a eukaryotic promoter and a DNA sequence which encodes a truncated VRP (or a functional derivative thereof) does not contain any intervening codons which are capable of encoding a methionine (i.e., AUG). The presence of such codons results either in a formation of a fusion protein (if the AUG codon is in the same reading frame as the truncated VRP coding sequence) or a frame-shift mutation (if the AUG codon is not in the same reading frame as the truncated VRP coding sequence).

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A truncated VRP nucleic acid molecule and an operably linked promoter may be introduced into a recipient prokaryotic or eukaryotic cell either as a nonreplicating DNA (or RNA) molecule, which may either be a linear molecule or, more preferably, a closed covalent circular molecule. Since such molecules are incapable of autonomous replication, the expression of the gene may occur through the transient expression of the introduced sequence. Alternatively, permanent expression may occur through the integration of the introduced DNA sequence into the host chromosome.

A vector may be employed which is capable of integrating the desired gene sequences into the host cell chromosome. Cells which have stably integrated the introduced DNA into their chromosomes can be selected by also introducing one or more markers which allow for selection of host cells which contain the expression vector. The marker may provide for prototrophy to an auxotrophic host, biocide resistance, e.g., antibiotics, or heavy metals, such as copper, or the like. The selectable

y (8 m.) (1 m.)

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needed for optimal synthesis of single chain binding protein mRNA. These elements may include splice signals, as well as transcription promoters, enhancers, and termination signals. cDNA expression vectors incorporating such elements include those described by Okayama, Molec. Cell. Biol. 3:280 (1983).

The introduced nucleic acid molecule can be incorporated into a plasmid or viral vector capable of autonomous replication in the recipient host. Any of a wide variety of vectors may be employed for this purpose. Factors of importance in selecting a particular plasmid or viral vector include: the ease with which recipient cells that contain the vector may be recognized and selected from those recipient cells which do not contain the vector; the number of copies of the vector which are desired in a particular host; and whether it is desirable to be able to "shuttle" the vector between host cells of different species.

Preferred eukaryotic plasmids include, for example, BPV, vaccinia, SV40, 2-micron circle, and the like, or their derivatives. Such plasmids are well known in the art (Botstein et al., Miami Wntr. Symp. 19:265-274(1982); Broach, In: "The Molecular Biology of the Yeast Saccharomyces: Life Cycle and Inheritance", Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, p. 445-470 (1981); Broach, Cell 28:203-204 (1982); Bollon et al., J. Clin. Hematol. Oncol. 10:39-48 (1980); Maniatis, In: Cell Biology: A Comprehensive Treatise, Vol. 3, Gene Sequence Expression, Academic Press, NY, pp. 563-608(1980).

Once the vector or nucleic acid molecule containing the construct(s) has been prepared for expression, the DNA construct(s) may be introduced into an appropriate host cell by any of a variety of suitable means, i.e., transformation,

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transfection, and the like. The most effective method for transfection of eukaryotic cell lines with plasmid DNA varies with the given cell type. After the introduction of the vector, recipient cells are grown in a selective medium, which selects for the growth of vector-containing cells. Expression of the cloned gene molecule(s) results in the production of truncated VRP or fragments thereof. This can take place in the transformed cells as such, or following the induction of these cells to differentiate (for example, by administration of bromodeoxyuracil to neuroblastoma cells or the like). A variety of incubation conditions can be used to form the peptide of the present invention. The most preferred conditions are those which mimic physiological conditions.

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transfectants, Production of the stable may accomplished by, for example, transfection of an appropriate cell line with an eukaryotic expression vector, such as pCEP4, in which the coding sequence for the truncated VRP polypeptide or subunit has been cloned into the multiple cloning site. These expression vectors contain a promoter region, such as the human cytomegalovirus promoter (CMV), that drive high-level transcription of desired DNA molecules in a variety of mammalian cells. In addition, these vectors contain genes for the selection of cells that stably express the DNA molecule of interest. The selectable marker in the pCEP4 vector encodes an enzyme that confers resistance to hygromycin, a metabolic inhibitor that is added to the culture to kill nontransfected cells.

Cells that have stably incorporated the transfected DNA will be identified by their resistance to selection media, as described above, and clonal cell lines will be produced by expansion of resistant colonies. The expression of the

#### Pharmaceutical Compositions and Therapeutic Uses

One object of this invention is to provide truncated VRP in a pharmaceutical composition suitable for therapeutic use. Thus, in one aspect the invention provides a method for stimulating angiogenesis in a patient by administering a therapeutically effective amount of pharmaceutical composition comprising a truncated VRP.

By "therapeutically effective amount" is meant an amount of a compound which produces the desired therapeutic effect in a patient. For example, in reference to a disease or disorder, it is the amount which reduces to some extent one or more symptoms of the disease or disorder, and returns to normal, either partially or completely, physiological or biochemical parameters associated or causative of the disease or disorder.

When used to therapeutically treat a patient it is an amount expected to be between 0.1 mg/kg to 100 mg/kg, preferably less than 50 mg/kg, more preferably less than 10 mg/kg, more preferably less than 10 mg/kg, more preferably less than 2 mg/kg. The amount of compound depends on the age, size, and disease associated with the patient.

The optimal formulation and mode of administration of compounds of the present application to a patient depend on factors known in the art such as the particular disease or disorder, the desired effect, and the type of patient. While the compounds will typically be used to treat human patients, they may also be used to treat similar or identical diseases in other vertebrates such as other primates, farm animals such as swine, cattle and poultry, and sports animals and pets such as horses, dogs and cats.

Preferably, the therapeutically effective amount is provided as a pharmaceutical composition. A pharmacelogical agent or composition refers to an agent or composition in a

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injection. Such forms should allow the agent or composition to reach a target cell whether the target cell is present in a multicellular host or in culture. For example, pharmacological agents or compositions injected into the blood stream should be soluble. Other factors are known in the art, and include considerations such as toxicity and forms which prevent the agent or composition from exerting its effect.

The claimed compositions can also be formulated as pharmaceutically acceptable salts (e.g., acid addition salts) and/or complexes thereof. Pharmaceutically acceptable salts are non-toxic salts at the concentration at which they are administered. The preparation of such salts can facilitate the pharmacological use by altering the physical-chemical characteristics of the composition without preventing the composition from exerting its physiological effect. Examples of useful alterations in physical properties include lowering the melting point to facilitate transmucosal administration and increasing the solubility to facilitate the administration of higher concentrations of the drug.

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Pharmaceutically acceptable salts include acid addition 20 salts such as those containing sulfate, hydrochloride, phosphate, sulfonate, sulfamate, sulfate, acetate, citrate, lactate, tartrate, methanesulfonate, ethanesulfonate, benzenesulfonate, p-toluenesulfonate, cyclolexylsulfonate, cyclohexylsulfamate and quinate. Pharmaceutically acceptable 25 salts can be obtained from acids such as hydrochloric acid, sulfuric acid, phosphoric acid, sulfonic acid, sulfamic acid, acetic acid, citric acid, lactic acid, tartaric acid, malonic acid, methanesulfonic acid, ethanesulfonic 30 benzenesulfonic acid, p-toluenesulfonic cyclcohexylsulfonic acid, cyclohexylsulfamic acid, and quinic

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medium in which the salt is insoluble, or in a solvent such as water which is then removed in vacuo or by freeze-drying or by exchanging the ions of an existing salt for another ion on a suitable ion exchange resin.

Carriers or excipients can also be used to facilitate administration of the compound. Examples of carriers and excipients include calcium carbonate, calcium phosphate, various sugars such as lactose, glucose, or sucrose, or types of starch, cellulose derivatives, gelatin, vegetable oils, polyethylene glycols and physiologically compatible solvents. The compositions or pharmaceutical composition can be administered by different routes including intravenously, intraperitoneal, subcutaneous, and intramuscular, orally, topically, or transmucosally.

The desired isotonicity may be accomplished using sodium chloride or other pharmaceutically acceptable agents such as dextrose, boric acid, sodium tartrate, propylene glycol, polyols (such as mannitol and sorbitol), or other inorganic or organic solutes. Sodium chloride is preferred particularly for buffers containing sodium ions.

The compounds of the invention can be formulated for a variety of modes of administration, including systemic and localized administration. Techniques topical or generally may be formulations found in Remington's Pharmaceutical Sciences, 18th Edition, Mack Publishing Co., Easton, PA, 1990. See also Wang, Y.J. and Hanson, M.A. "Parenteral Formulations of Proteins and Peptides: Stability and Stabilizers," Journal of Parenteral Science and Technology, Technical Report No. 10, Supp. 42:2S (1988). A suitable administration format may best be determined by a medical practitioner for each patient individually.

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compounds of the invention are formulated in liquid solutions, preferably in physiologically compatible buffers such as Hank's solution or Ringer's solution. Alternatively, the compounds of the invention are formulated in one or more excipients (e.g., propylene glycol) that are generally accepted as safe as defined by USP standards. They can, for example, be suspended in an inert oil, suitably a vegetable oil such as sesame, peanut, olive oil, or other acceptable carrier. Preferably, they are suspended in an aqueous carrier, for example, in an isotonic buffer solution at a pH of about 5.6 to 7.4. These 10 compositions may be sterilized by conventional sterilization techniques, or may be sterile filtered. The compositions may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions, such as pH buffering agents. Useful buffers include for example, sodium 15 acetate/acetic acid buffers. A form of repository or "depot" slow release preparation may be used so that therapeutically effective amounts of the preparation are delivered into the bloodstream over many hours or days following transdermal injection or delivery. In addition, the compounds may be 20 and redissolved or suspended formulated in solid form immediately prior to use. Lyophilized forms are also included.

Systemic administration can also be by transmucosal or transdermal means, or the molecules can be administered orally.

25 For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art, and include, for example, for transmucosal administration, bile salts and fusidic acid derivatives. In addition, detergents

30 may be used to facilitate permeation. Transmucosal administration may be, for example, through nasal sprays or

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For topical administration, the compounds of the invention are formulated into ointments, salves, gels, or creams, as is generally known in the art.

If desired, solutions of the above compositions may be thickened with a thickening agent such as methyl cellulose. They may be prepared in emulsified form, either water in oil or oil in water. Any of a wide variety of pharmaceutically acceptable emulsifying agents may be employed including, for example, acacia powder, a non-ionic surfactant (such as a Tween), or an ionic surfactant (such as alkali polyether alcohol sulfates or sulfonates, e.g., a Triton).

Compositions useful in the invention are prepared by mixing the ingredients following generally accepted procedures. For example, the selected components may be simply mixed in a blender or other standard device to produce a concentrated mixture which may then be adjusted to the final concentration and viscosity by the addition of water or thickening agent and possibly a buffer to control pH or an additional solute to control tonicity.

The amounts of various compounds of this invention to be administered can be determined by standard procedures. Generally, a therapeutically effective amount is between about 1 nmole and 3 µmole of the molecule, preferably between about 10 nmole and 1 µmole depending on the age and size of the patient, and the disease or disorder associated with the patient. Generally, it is an amount between about 0.1 and 50 mg/kg, preferably 1 and 20 mg/kg of the animal to be treated.

For use by the physician, the compositions will be provided in dosage unit form containing an amount of a truncated VRP, VRP polypeptide, or VRP subunit.

(1992)). Miller states that advances have resulted in practical approaches to human gene therapy that have demonstrated positive initial results. The basic science of gene therapy is described in Mulligan, Science 260:926-931 (1993). One example of gene therapy is presented in Example 7, which describes the use of adenovirus-mediated gene therapy.

As another example, an expression vector containing the truncated VRP coding sequence may be inserted into cells, the cells are grown in vitro and then injected or infused in large numbers into patients. In another example, a DNA segment containing a promoter of choice (for example a strong promoter) is transferred into cells containing an endogenous truncated VRP in such a manner that the promoter segment enhances expression of the endogenous truncated VRP gene (for example, the promoter segment is transferred to the cell such that it becomes directly linked to the endogenous truncated VRP gene).

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The gene therapy may involve the use of an adenovirus vector including a nucleotide sequence coding for a truncated VRP subunit, or a naked nucleic acid molecule coding for a truncated VRP subunit. Alternatively, engineered cells containing a nucleic acid molecule coding for a truncated VRP subunit may be injected. Example 7 illustrates a method of gene therapy using an adenovirus vector to provide angiogenesis therapy.

Expression vectors derived from viruses such as retroviruses, vaccinia virus, adenovirus, adeno-associated virus, herpes viruses, several RNA viruses, or bovine papilloma virus, may be used for delivery of nucleotide sequences (e.g., cDNA) encoding recombinant truncated VRP subunit into the targeted cell population. Methods which are well known to those skilled in the art can be used to construct recombinant

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(1989), and in Ausubel et al., <u>Current Protocols in Molecular Biology</u>, Greene Publishing Associates and Wiley Interscience, N.Y. (1989). Alternatively, recombinant nucleic acid molecules encoding protein sequences can be used as naked DNA or in reconstituted system <u>e.g.</u>, liposomes or other lipid systems for delivery to target cells (<u>See e.g.</u>, Felgner et al., Nature 337:387-8, 1989). Several other methods for the direct transfer of plasmid DNA into cells exist for use in human gene therapy and involve targeting the DNA to receptors on cells by complexing the plasmid DNA to proteins. <u>See</u>, Miller, Nature 357:455-60, 1992.

In its simplest form, gene transfer can be performed by simply injecting minute amounts of DNA into the nucleus of a cell, through a process of microinjection. Capecchi MR, Cell 22:479-88 (1980). Once recombinant genes are introduced into a 15 cell, they can be recognized by the cells normal mechanisms for transcription and translation, and a gene product will be Other methods have also been attempted for introducing DNA into larger numbers of cells. These methods transfection, wherein DNA is precipitated with 20 calcium phosphate and taken into cells by pinocytosis (Chen C. 7:2745-52 Mol. Cell Biol. (1987));Okayama H, and electroporation, wherein cells are exposed to large voltage pulses to introduce holes into the membrane (Chu G. et al., Nucleic Acids Res., 15:1311-26 (1987)); lipofection/liposome 25 fusion, wherein DNA is packaged into lipophilic vesicles which fuse with a target cell (Felgner PL., et al., Proc. Natl. Acad. Sci. USA. 84:7413-7 (1987)); and particle bombardment using DNA bound to small projectiles (Yang NS. et al., Proc. Natl. Acad. Sci. 87:9568-72 (1990)). Another method for introducing DNA into cells is to couple the DNA to chemically DNA into cells. The admixture of adenovirus to solutions containing DNA complexes, or the binding of DNA to polylysine covalently attached to adenovirus using protein crosslinking agents substantially improves the uptake and expression of the recombinant gene. Curiel DT et al., Am. J. Respir. Cell. Mol. Biol., 6:247-52 (1992).

In addition, it has been shown that adeno-associated virus vectors may be used for gene delivery into vascular cells (Gnatenko, D., J. of Invest. Med. 45:87-97, (1997)).

As used herein "gene transfer" means the process of 10 introducing a foreign nucleic acid molecule into a cell. Gene transfer is commonly performed to enable the expression of a particular product encoded by the gene. The product may include a protein, polypeptide, anti-sense DNA or RNA, or enzymatically active RNA. Gene transfer can be performed in cultured cells or by direct administration into animals. Generally gene transfer involves the process of nucleic acid molecule contact with a target cell by non-specific or receptor mediated interactions, uptake of nucleic acid molecule into the cell through the membrane or by endocytosis, and release of 20 nucleic acid molecule into the cytoplasm from the plasma membrane or endosome. Expression may require, in addition, movement of the nucleic acid molecule into the nucleus of the binding to appropriate nuclear factors for cell and 25 transcription.

As used herein "gene therapy" is a form of gene transfer and is included within the definition of gene transfer as used herein and specifically refers to gene transfer to express a therapeutic product from a cell in vivo or in vitro. Gene transfer can be performed ex vivo on cells which are then transplanted into a patient, or can be performed by direct

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In another preferred embodiment, a vector having nucleic acid molecule sequences encoding a truncated VRP is provided in which the nucleic acid molecule sequence is expressed only in a specific tissue. Methods of achieving tissue-specific gene expression as set forth in International Publication No. WO 93/09236, filed November 3, 1992 and published May 13, 1993.

In another preferred embodiment, a method of gene replacement is set forth. "Gene replacement" as used herein means supplying a nucleic acid molecule sequence which is capable of being expressed in vivo in an animal and thereby providing or augmenting the function of an endogenous gene which is missing or defective in the animal.

In all of the preceding vectors set forth above, a further aspect of the invention is that the nucleic acid sequence contained in the vector may include additions, deletions or modifications to some or all of the sequence of the nucleic acid, as defined above.

#### Examples

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To assist in understanding the present invention, the following Examples are included which describes the results of a series of experiments. The experiments relating to this invention should not, of course, be construed as specifically limiting the invention and such variations of the invention, now known or later developed, which would be within the purview of one skilled in the art are considered to fall within the scope of the invention as described herein and hereinafter claimed.

### Example 1

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Cloning of N-Terminally Truncated VEGF-B, (des-(1-20)-p21-VEGF-B (or des(2-21)-VEGF-B).

In order to create a novel VEGF-B-related protein that lacks the first 20 amino acids, a cDNA construct is created in the following manner:

A DNA encoding human VEGF-B is amplified from a human heart or skeletal muscle cDNA), or a human fetal brain cDNA library, or a cDNA preparation from another suitable human tissue source by PCR with oligonucleotides corresponding to the published sequence of human VEGF-B. Using standard molecular biology techniques (Sambrook et al., Molecular Cloning, A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor NY), a DNA fragment then is generated that encodes at its 5' end the signal sequence of human VEGF-B, followed by a codon for proline, the first amino acid residue in mature VEGF-B, and then followed by codons corresponding to amino acids from residues 22 to the C-terminus of human VEGF-B, followed by a stop codon. Appropriate additional non-coding nucleotide sequences are added to the 5' and 3' ends of this DNA construct so as to allow insertion of the DNA into an appropriate expression vector.

In this manner the cleavage site for the signal peptide is preserved in a manner identical to that found in native VEGF-B. However, this strategy results in a change in the new N-terminal amino acid of the truncated VEGF-B. Whereas the normal N-terminal amino acid residue in des(1-20)-VEGF-B is a tyrosine residue:

mspllrrlllvallqlartqa[PVSQFDGPSHQKKVVPWIDV] YTRAT, the new N-terminal amino acid is proline, and the resulting truncated VEGF-B is equivalent to des(2-21)-VEGF-B):

not expected to have any effect on the biological activity of the truncated VEGF-B. The advantage of this strategy is that the signal peptide sequence is maintained thus ensuring efficient cleavage of the signal peptide from the precursor during protein processing/secretion.

In another example, truncated VEGF-B, des(1-15)-VEGF-B, is constructed by deleting the first 15 amino acids. The signal peptide cleavage site would be preserved in this case because residue#16 and residue#1 (the new and old N-termini) are identical (proline):

mspllrrillvallqlartqa[PVSQFDGPSHQKKVV]PWIDVYTRAT...

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mspllrrillvallqlartqaPWIDVYTRAT..

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One of skill in the art would understand that other signal peptides may be used in the present invention. For example, the signal peptide of VEGF-B or VEGF-C could be used which would require that the first amino acid of the truncated protein be an alanine or glycine, respectively, in order to preserve the respective signal peptide cleavage sites. A further alternative would be to use signal peptide sequences from other known proteins; some of these may have cleavage sites compatible with the N-terminal tyrosine of the truncated des(1-20)-VEGF-B.

Another alternative would be to generate a construct that encodes a precursor protein with a cleavage site that incorporates two, rather than one, amino acids from the N-terminus of the original VEGF-B protein sequence. The purpose of this strategy would be to ensure more fully that the cleavage site is compatible with signal peptidase function.

This would introduce two new amino acids at the N-terminus of the truncated VEGF-B sequence but such a change would not be expected.

The strategy described to generate DNA for expression of des(1-20)-VEGF-B is useful for generation in an analogous manner of VEGF-B mutants with N-terminal truncations of other desired lengths. Further, the strategy is useful to generate N-terminal truncations of other desired lengths in other VEGF-related forms and their isoforms of other species.

# Example 2: Expression Of N-Terminally Truncated VEGF-B Subunits

The DNA fragment encoding truncated VEGF-B from Example 1 10 may be cloned into a suitable plasmid vector.

Sf9 (Sporoptera frugiperda) cells are co-transfected with baculovirus transfer vector pAcUW51 containing cDNA encoding truncated VEGF-B and baculovirus (Baculogold, Pharmingen, San Diego, CA). Selection and plaque purification of recombinant virus are performed according to established protocols using 15 Blue agar overlays (Gibco BRL). High stock of recombinant virus is produced in exponentially growing Sf9 cells using a multiplicity of infection of 0.05. For expression of truncated VEGF-B, Sf9 cells (1x106 cells/ml) growing in serum free medium 20 are infected with recombinant virus at a multiplicity of 10. Supernatant is collected after 72 hours post infection. VEGF expression in baculovirus-infected insect cells, which can be used to express the truncated VRPs of the present invention is also described in Fiebich et al., (Eur. J. Biochem. 211: 19-26, 1993). In this system, VEGF has been shown to be produced in 2.5 high yield, with efficient glycosylation similar to that seen in mammalian cells. In fact, those skilled in the art will recognize that expression in other systems, including mammalian cell expression systems, is considered to be within the scope of this invention. Methods of expressing VEGF proteins which 30 can be used to express the truncated VRPs of the present

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(Baculovirus Expression Vectors: A Laboratory Manual (W.H. Freeman, New York), 1992).

Those skilled in the art will recognize that other expression systems may also be used to express functionally active truncated VRPs.

Functionally active recombinant VEGF isoforms have been expressed in E. Coli (Wilting et al., <u>Dev. Biol.</u> 176, 76-85, 1996) from inclusion body by refolding according to the procedure described previously for homo- and heterodimers of PDGF (Schneppe et al., Gene 143, 201-09, 1994) and in yeast (Mohanraj et al., <u>Biochem. Biophys. Res. Commun.</u> 215:750-56, 1995).

Still other methods of expressing VEGF which can be used to express VRPs in the present invention are described, for example, in Jasny, <u>Science</u> 238:1653, 1987; and Miller et al., In: Genetic Engineering, 1986), Setlow, J.K., et al., eds., Plenum, Vol. 8, pp. 277-297).

#### Example 3: Purification Of Recombinant Truncated VRPS

Por purification of the baculovirus-expressed truncated VEGF-B of Example 2 from insect-cell supernatant, a number of standard techniques can be used. These techniques include, but are not limited to ammonium sulfate precipitation, acetone precipitation, ion-exchange chromatography, size exclusion chromatography, hydrophobic interaction chromatography, reverse-phase HPLC, concanavalin A affinity chromatography, isoelectric focusing, and chromatofocusing. Other standard protein purification techniques are readily obvious to one skilled in the art. For example, proteins with specific tags, such as histidine tags, antigen tags, etc., could be produced by engineering DNA encoding such tags into the VEGF-B DNA such

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methods are considered within the scope of the present invention.

A preferred purification method for truncated forms of VEGF-B is described in the following: Sf9 Cell supernatant is centrifuged at 10000 rpm for 30 minutes to remove cell debris and viral particles. Supernatant is then concentrated and dialyzed against 20 mM Tris (pH 8.3) for 24 hours. dialyzed supernatant is centrifuged again to remove insoluble material and loaded onto a Sepharose Q anion exchange column. Protein is eluted from the column by gradient elution using a 10 gradient of NaCl (0 - 1 M NaCl). Chromatography fractions are analyzed by SDS polyacrylamide gel electrophoresis and by ELISA using an antibody that recognizes VEGF-B. Fractions with VEGF-B immunoreactivity are pooled, concentrated, and dialyzed overnight against 0.1% trifluoroacetic acid. Material so 15 prepared is further purified by reverse phase HPLC. Typically approximately 2-5 mg of protein is loaded on a semipreparative C4 column and eluted with a gradient of acetonitrile in 0.1% trifluoracetic acid as described in Esch et al., Meth. Enzymol. 103, 72-89, 1983. Fractions containing truncated VEGF-B are 20 pooled and stored at -80 degrees Celsius until further use.

A preferred method of purification of the basic and heparin-binding N-terminally truncated forms of VEGF-related protein subunits and analogs thereof includes the combined use of heparin-Sepharose affinity chromatography and cation-exchange chromatography, optionally followed by reverse-phase HPLC, essentially as described in Connolly et al., <u>J. Biol. Chem.</u> 264:20017-24, 1989, Gospodarowicz et al., (<u>Proc. Natl. Acad. Sci. USA</u>, 86:7311-15, 1989), or Plouet et al., (<u>Embo J. 8:3801-06</u>, 1989).

Purification is monitored by following the elution of VRP-

preparations consisting of cells or cell membrane preparations in functional assays as described in Examples 4-6.

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The truncated VRPs expressed in other eukaryotic cell systems such as yeast or mammalian cells, may be purified in the same manner.

Truncated VRPs expressed in prokaryotic cells will likely need to undergo a re-folding step for proper dimerization of subunits, as described in, for example, Schneppe et al., (Gene 143:201-09, 1994).

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#### Example 4: Receptor-Binding Assay

The binding of truncated VRPs to VEGF receptors can be assessed in various ways. Useful methods include the determination of the ability of VRP analogs to bind to endothelial cells or to cells artificially transfected with KDR, or to soluble forms of the KDR receptor (for example, a KDR/alkaline phosphatase fusion protein (Gitay-Goren et al., J. Biol. Chem. 271:5519-23 (1996)). A preferred procedure has been described by Terman et al. (Biochem. Biophys. Res. Commun. 187:1579-86, 1992).

In this procedure, KDR cDNA is transfected into CMT-3 monkey kidney cells by the DEAE-dextran method by incubating plated cells with DMEM containing 1  $\mu$ g/ml DNA, 0.5  $\mu$ g/ml DEAE dextran, and 100  $\mu$ M chloroquine. Following incubation for 4 hours at 37 degrees Celsius, the medium is aspirated and cells are exposed to 10% DMSO in PBS for one minute. The cells are then washed once with DMEM containing 10% calf serum and then incubated for 40 hours at 37 degrees Celsius in DMEM/10% calf serum containing 100  $\mu$ M ZnCl<sub>2</sub> and 1  $\mu$ M CdCl<sub>2</sub>.

30 · VEGF-B is radiolodinated using either the Iodogen method or the chloramine T method. Padiolabelled VEGE-B is constrated

order of  $10^5$  cpm/ng. For radioceptor assays, CMT-3 ( $10^5$  cells/well) are plated in 12-well plates. Twenty four hours later, cells are washed twice with PBS, and 0.5 ml of DMEM containing 0.15% gelatin and 25 mM HEPES, pH 7.4 is added.  $^{125}\text{I-VEGF-B}$ , at concentrations ranging from 1-500 pM, is then added. Binding experiments are done in the presence or absence of 0.5 nM unlabeled VEGF-B for the determination of specific binding. After a 90-minute incubation at room temperature, a 50  $\mu$ l sample of the media from each well is used to determine the concentration of free radioligand, and the wells are washed 3 times with ice cold PBS containing 0.1% BSA. Cells are extracted from the wells by incubation for 30 minutes with 1% Triton X100 in 100 mM sodium phosphate, pH 8.0, and the radioactivity of the extract is determined in a gamma counter.

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#### Example 5: Mitogenic Assay

The mitogenic activity of truncated VRPs on endothelial cells of human or mammalian origin can be determined by a number of different procedures, including assays where cell proliferation is measured by growth of cell numbers or by incorporation of radioactive DNA precursors (thymidine incorporation) or otherwise appropriately labeled precursors (bromo-deoxyuridine incorporation). These and other methods generally used to determine cell proliferation, including those methods where mitogenic activity is assessed in vivo (for example by determining the mitotix index of endothelial cells) are considered within the scope of this invention. A preferred method is described herein (Bohlen et al., Proc. Natl. Acad. Sci. USA 81:5364-68, 1994): aortic arch endothelial cells maintained in stock cultures in the presence of Dulbecco's modified Eagle's medium supplemented

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ratio of 1:64. For mitogenic assays, cell monolayers from stock plates (at passages 3-10) are dissociated using trypsin. Cells are then seeded at a density of approximately 8000 cells/well in 24-well plates in the presence of DMEM and antibiotics as described above. Samples to be assayed (1-10 μl), appropriately diluted in DMEM/0.1% bovine serum albumin), are added six hours after plating of cells and again after 48 hours. After 4 days of culture, endothelial cells are detached from plates with trypsin and counted using a Coulter particle counter.

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Another mitogenic activity assay is provided in Olofsson, B. et al., <u>Proc. Natl. Acad. Sci. USA</u> 93:2576-81, 1996). Second passage human umbilical vein endothelial cells (HUVECs) are plated into 96-well plates (4 X 103 cells per well) in M-199 medium supplemented with 10% (vol/vol) fetal bovine serum and incubated for 24 hours. Cell culture conditioned medium containing the truncated VRP, in the presence of 1-10  $\mu$ g/ml heparin, or purified truncated VRP is added to the HUVECs, and the cells are stimulated for 48 hours. Fresh cell culture conditioned medium containing [3H] thymidine (Amersham;  $\mu$ Ci/ml) is added to the cells and stimulation is continued for another 48 hours. Cells are washed with PBS and trypsinized and the incorporated radioactivity is determined by liquid scintillation counting. The activity of truncated VRP is compared to the activity of non-truncated VRP.

another alternative method, bovine capillary endothelial (BCE) cells are seeded into 24-well plates and grown until confluence in minimal essential medium (MEM) supplemented with 10% fetal calf serum. Cells are starved in MEM supplemented with 3% fetal calf serum for 72 hours, after which conditioned medium diluted into serum-free medium is

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with NaOH, and incorporated radioactivity is determined by liquid scintillation counting. The activity of truncated VRP is compared to that of non-truncated VRP. Bovine fibroblast growth factor (b-FGF) may be used as an additional control for mitogenic activity, and may also be used to measure its potentiating activity of truncated VRP activity.

## Example 6: Angiogenic Activity Of Truncated VRPS

The angiogenic activity of substances can be determined using a variety of in vivo methods. Commonly used methods include the chick chorioallantoic membrane assay, the corneal pouch assay in rabbits, rats, or mice, the matrigel implant assay in mice, the rabbit ear chamber angiogenesis assay, the hamster cheek pouch assay, the Hunt-Schilling chamber model and the rat sponge implant model. Other assay methods to assess the formation of new blood vessels have been described in the literature and are considered to be within the scope of this invention.

A preferred method for demonstrating the angiogenic activity of truncated VRPs is the rabbit corneal pouch assay. In this assay, Elvax (ethylene vinyl acetate) polymer pellets containing approximately 1-1000 ng of the growth factor and a constant amount of rabbit serum albumin as carrier is implanted into a surgical incision in the cornea as described in more detail in Phillips and Knighton, Wound Rep. Reg. 3, 533-539, 1995; Gimbrone et al., J. Natl. Canc. Inst. 52:413-27, 1974; Risau, Proc. Natl. Acad. Sci. USA 83:3855-59, 1986). Growth factor-induced vascularization of the cornea is then observed over a period of 2 weeks. Semiquantitative analysis is possible with morphometric and image analysis techniques using photographs of corneas.

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# Example 7: Gene-Transfer-Mediated Angiogenesis Therapy Using Truncated VRPS

Truncated VRPs are used for gene-transfer-mediated angiogenesis therapy as described, for example, in PCT/US96/02631, published September 6, 1996 as WO96/26742, hereby incorporated by reference herein in its entirety.

#### Adenoviral Constructs

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helper independent replication deficient human Α adenovirus 5 system may be used for gene-transfer. A nucleic acid molecule coding for a truncated VRP subunit may be cloned into the polylinker of plasmid ACCMVPLPA which contains the CMV promoter and SV40 polyadenylation signal flanked by partial adenoviral sequences from which the E1A and (essential for viral replication) have been deleted. plasmid is co-transferred (lipofection) into 293 cells with plasmid JM17 which contains the entire human adenoviral 5 genome with an additional 4.3 kb insert making pJM17 too large to be encapsidated. Homologous rescue recombination results in adenoviral vectors containing the transgene in the absence of Although these recombinants E1A/E1B sequences. nonreplicative in mammalian cells, they can propagate in 293 cells which have been transformed with E1A/E1B and provided these essential gene products in trans. Transfected cells are monitored for evidence of cytopathic effect which usually occurs 10-14 days after transfection. To identify successful recombinants, cell supernatant from plates showing a cytopathic effect is treated with proteinase K (50 mg/ml with 0.5% sodium dodecyl sulfate and 20 mM EDTA) at 56°C for 60 minutes, phenol/

promoter and SV40 polyadenylation sequences to amplify the truncated VRP subunit nucleic acid insert and (Biotechniques 15:868-72, 1993) designed to concomitantly amplify adenoviral sequences. Successful recombinants then are plaque purified twice. Viral stocks are propagated in 293 cells to titers ranging between  $10^{10}$  and  $10^{12}$  viral particles, and are purified by double CsCl gradient centrifugation prior to use. The system used to generate recombinant adenoviruses imposed a packing limit of 5kb for transgene inserts. truncated VRP genes, driven by the CMV promoter and with the 10 SV40 polyadenylation sequences are well within the packaging Recombinant vectors are plaque purified by constraints. The resulting viral vectors standard procedures. are propagated on 293 cells to titers in the  $10^{10}-10^{12}$ particles range. Cells are infected at 80% confluence and 15 harvested at 36-48 hours. After freeze-thaw cycles the cellular debris is pelleted by standard centrifugation and the further purified by CsCl double ultracentrifugation (discontinuous 1.33/1.45 CsCl gradient; cesium prepared in 5 mM Tris, 1 mM EDTA (pH 7.8); 90,000 x g (2 20 hr), 105,000 x g (18 hr)). Prior to in vivo injection, the viral stocks are desalted by gel filtration through Sepharose columns such as G25 Sephadex. The resulting viral stock has a final viral titer approximately in the  $10^{10}\text{--}10^{12}$  viral particles range. The adenoviral construct should thus be highly 25 purified, with no wild-type (potentially replicative) virus.

#### Porcine Ischemia Model For Angiogenesis

A left thoracotomy is performed on domestic pigs (30-40 kg) under sterile conditions for instrumentation. (Hammond, et al. J Clin Invest. 92:2644-52 (1993); Roth, et al. J. Clin.

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atrium to permit ECG recording and atrial pacing. Finally, an ameroid constrictor (ameroid), a metal ring including an ameroid substance, is placed around the proximal left circumflex coronary artery (LCx) (Hammond et al. J. Clin. Invest. 92:2644-52 (1993)). After a stable degree of ischemia develops, the treatment group receives an adenoviral construct that includes a truncated VRP gene driven by a CMV promoter. Control animals receive gene transfer with an adenoviral construct that includes a reporter gene, lacZ, driven by a CMV promoter.

Studies are initiated 35 + 3 days after ameroid placement, at a time when collateral vessel development and pacing-induced dysfunction are stable (Roth, et al. Am J Physiol 253:1-11279et al. Circulation 82:1778-89). Roth, 1288, 1987, and Conscious animals are suspended in a sling and pressures from 15 the left ventricle (LV), left atrium (LA) and aorta, and electrocardiogram are recorded in digital format on-line (at rest and during atrial pacing at 200 bpm). Two-dimensional and M-mode images are obtained using a Hewlett Packard ultrasound imaging system. Images are obtained from a right parasternal 20 approach at the mid-papillary muscle level and recorded on VHS tape. Images are recorded with animals in a basal state and again during right atrial pacing (HR=200 bpm). These studies are performed one day prior to gene transfer and repeated 14 + 25 1 days later. Rate-pressure products and left atrial pressures should be similar in both groups before and after gene transfer, indicating similar myocardial oxygen demands and loading conditions. Echocardiographic measurements are made using standardized criteria (Sahn, et al. Circulation 58:1072, 1978). End-diastolic wall thickness (EDWTh) and end-systolic 30 wall thickness (ESWTh) are measured from 5 continuous beats and

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demonstrate reproducibility of echocardiographic measurements, animals should be imaged on two consecutive days, showing high correlation ( $r^2=0.90$ ; p=0.005).

35 ± 3 days after ameroid placement, well after ameroid closure, but before gene transfer, contrast echocardiographic studies are performed using the contrast material (Levovist) which is injected into the left atrium during atrial pacing (200 bprn). Studies are repeated 14 ± 1 days after gene transfer. Peak contrast intensity is measured from the video images using a computer-based video analysis program (Color Vue II, Nova Microsonics, Indianapolis, Indiana), that provides an objective measure of video intensity. The contrast studies are analyzed without knowledge of which gene the animals have received.

15 At completion of the study, animals are anesthetized and midline thoracotomy performed. The brachycephalic artery is isolated, a canula inserted, and other great vessels ligated. The animals receive intravenous heparin (10,000 IU) papaverine (60 mg). Potassium chloride is given to induce diastolic cardiac arrest, and the aorta cross-clamped. Saline 20 is delivered through the brachycephalic artery cannula (120 mmHg pressure), thereby perfusing the coronary arteries. Glutaraldehyde solution (6.25%, 0.1 M cacodylate buffer) was perfused (120 mmH pressure) until the heart is well fixed (10-25 15 min). The heart is then removed, the beds identified using color-coded dyes injected anterograde through the left anterior descending (LAD), left circumflex (LCx), and right coronary arteries. The ameroid is examined to confirm closure. Samples taken from the normally perfused and ischemic regions are 30 divided into thirds and the endocardial and epicardial thirds are plastic-imbedded. Microscopic analysis to quantitate

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and epicardium of each region) and point-counting is used to determine capillary number per fiber number ratio at 400X magnification. Twenty to twenty-five high power fields are counted per subsample. Within each region, capillary number to fiber number rations should be similar in endocardium and epicardium so the 40-50 field per region should be averaged to provide the transmural capillary to fiber number ratio.

To establish that improved regional function and blood flow result from transgene expression, PCR and RT-PCR may be used to detect transgenic truncated VRP DNA and mRNA in myocardium from animals that have received truncated VRP gene transfer. Using a sense primer to the CMV promoter [GCAGAGCTCGTTTAGTGAAC] [SEQ I.D. NO. 41]; and an antisense primer to the internal truncated VRP subunit sequence, PCR is used to amplify the expected 500 bp fragment. Using a sense primer to the beginning of the truncated VRP subunit sequence, and an antisense primer to the internal truncated VRP sequence, RT-PCR is used to amplify the expected 400 bp fragment.

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Finally, using a polyclonal antibody directed against VRP, truncated VRP expression may be demonstrated 48 hours as well as  $14 \pm 1$  days after gene transfer in cells and myocardium from animals that have received gene transfer with a truncated VRP gene.

The helper independent replication deficient human adenovirus 5 system is used to prepare transgene containing vectors. The material injected in vivo should be highly purified and contain no wild-type (replication competent) adenovirus. Thus adenoviral infection and inflammatory infiltration in the heart are minimized. By injecting the material directly into the lumen of the coronary artery by coronary catheters, it is possible to target the gene

be found in the urine at any time after intracoronary injection.

Injection of the construct (4.0 ml containing about 10<sup>11</sup> viral particles of adenovirus) is performed by injecting 2.0 ml into both the left and right coronary arteries (collateral flow to the LCx bed appeared to come from both vessels). Animals are anesthetized, and arterial access acquired via the right carotid by cut-down; a 5F Cordis sheath is then placed. A 5F Multipurpose (A2) coronary catheter is used to engage the coronary arteries. Closure of the LCx ameroid is confirmed by contrast injection into the left main coronary artery. The catheter tip is then placed 1 cm within the arterial lumen so that minimal material is lost to the proximal aorta during injection. This procedure is carried out for each of the pigs.

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Once gene transfer is performed, three strategies are used to establish successful incorporation and expression of the gene. (1) Some constructs may include a reporter gene (lacZ); (2) myocardium from the relevant beds is sampled, and immunoblotting is performed to quantitate the presence of truncated VRP and (3) PCR is used to detect truncated VRP mRNA and DNA.

The regional contractile function data obtained should show that control pigs show a similar degree of pacing-induced dysfunction in the ischemic region before and 14 ± 1 days after gene transfer. In contrast, pigs receiving truncated gene transfer should show an increase in wall thickening in the ischemic region during pacing, demonstrating that truncated VRP subunit gene transfer in accordance with the invention is associated with improved contraction in the ischemic region during pacing. Wall thickening in the normally perfused region (the interventricular septum) should be normal during pacing

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sonomicrometry during atrial pacing in the same model (Hammond, et al. <u>J. Clin. Invest.</u> 92:2644, 1993), documenting the accuracy of echocardiography for the evaluation of ischemic dysfunction.

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## Sequence Listing

	(1) GENE	RAL INFORMATION:	
5	(i)	APPLICANT:	Collateral Therapeutics
	(ii)	TITLE OF INVENTION:	TRUNCATED VEGF-RELATED PROTEINS
10	(iii)	NUMBER OF SEQUENCES:	41
-0	(iv)	CORRESPONDENCE ADDRESS:	
15		(A) ADDRESSEE: (B) STREET:	Lyon & Lyon 633 West Fifth Street Suite 4700
		(C) CITY: (D) STATE: (E) COUNTRY: (F) ZIP:	Los Angeles California U.S.A. 90071-2066
20	(v)		
		(A) MEDIUM TYPE:	3.5" Diskette, 1.44 Mb storage
25		(B) COMPUTER: (C) OPERATING SYSTEM: (D) SOFTWARE:	IBM Compatible IBM P.C. DOS 5.0 FastSEQ for Windows 2.0
30	(vi)	CURRENT APPLICATION DATA:	
		(A) APPLICATION NUMBER: (B) FILING DATE: (C) CLASSIFICATION:	08/842,984 April 25, 1997
35		(o, o <u>p.166 1</u> 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	
	(vii)	PRIOR APPLICATION DATA:	
40		(A) APPLICATION NUMBER: (B) FILING DATE:	
	(viii)	ATTORNEY/AGENT INFORMATION:	:
45		(A) NAME: (B) REGISTRATION NUMBER: (C) REFERENCE/EOCKET NUMBE	Warburg, Richard J. 32,327 ER: 221/062
50	(ix)	TELECOMMUNICATION INFORMATI	ION:
		(A) TELEPHONE: (B) TELEFAX: (C) TELEX:	(213) 489-1600 (213) 955-0440 67-3510
EE			

			(B)		PE: OPOLO	GY:				linea		La				
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10	Met 1	Ser	Pro	Leu	Leu 5	Arg	Arg	Leu	Leu	Leu 10	Val	Ala	Leu	Leu	Gln 15	Leu
	Ala	Arg	Thr	Gln 20	Ala	Pro	Val	Ser	Gln 25	Phe	Asp	Gly	Pro	Ser 30	His	Gln
15	Lys	Lys	Val 35	Val	Pro	Trp	Ile	Asp 40	Val	Тут	Thr	Arg	Ala 45	Thr	Cys	Gln
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20	Val 65	Lys	Gln	Leu	Val	Pro 70	Ser	Cys	Val	بدلين	Val 75	Gln	Arg	Сув	Cly	Cly 80
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35	Lys	Glu 130	Ser	Ala	Val	Lys	Pro 135	Asp	Ser	Pro	Arg	Ile 140	Leu	Cys	Pro	Pro
<i>J J</i>	Cys 145		Gln	Arg	Arg	Gln 150	Arg	Pro	Asp	Pro	Arg 155	Thr	Cys	Arg	Cys	Arg 160
40	Cys	Arg	Arg	Arg	Arg 165	Phe	Leu	His	Cys	Gln 170	Gly	Arg	Gly	Leu	Glu 175	Leu
	Asn	Pro	Asp	Thr 180	Cys	Arg	Cys	Arg	Lys 185		Arg	Lys				
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		(ii)	MO.	LECU	LE T	YPE:				Prot	ein					
55		(xi)	SEC	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	o:	2:				

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	Ala	Ala	Asp 195	Ala	Ala	Ala	Ser	Ser 200	Val	Ala	Lys	Gly	Gly 205	Ala		
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	Ala	Leu	Leu	Pro 20	Glγ	Pro	Arg	Glu	Ala 25	Pro	Ala	Ala	Ala	Ala 30	Ala	Phe
55	Glu	Ser	su Gly	Leu	Asp	Leu	Ser	Asp	Ala	Glu	Pro	Asp	Ala 1	Gly	Glu	Ala

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55	Leu	Λsn	Pro 355	Gly	Lys	Cys	Ala	Cys 361	Glu	Cys	Thr	Glu	Ser ngs	Pro	Gln	Lys

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Tyr	Ser	Glu	Glu	Val	Cys	Arg	Cys	Val	Pro	Ser	Tyr	Trp	Lys	Arg	Pro
				405					410					415	

- 5 Gln Met Ser
  - (2) INFORMATION FOR SEQ ID NO: 4:
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- (A) LENGTH:
- 170 amino acids

- amino acid
- (A) LENGTH:
  (B) TYPE:
  (D) TOPOLOGY:
- linear

- 15
- (ii) MOLECULE TYPE:

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
- 20 Met Pro Val Met Arg Leu Phe Pro Cys Phe Leu Gln Leu Leu Ala Gly
  - Leu Ala Leu Pro Ala Val Pro Pro Gln Gln Trp Ala Leu Ser Ala Gly
- 25 Asn Gly Ser Ser Glu Val Glu Val Val Pro Phe Gln Glu Val Trp Gly
- Arg Ser Tyr Cys Arg Pro Ile Glu Thr Leu Val Asp Ile Phe Gln Glu 30
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- 35 Met Arg Cys Gly Gly Cys Cys Asn Asp Glu Gly Leu Glu Cys Val Pro
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- Glu Cys Arg Pro Leu Arg Glu Lys Met Lys Pro Glu Arg Arg Arg Pro 45
  - Lys Gly Arg Gly Lys Arg Arg Glu Lys Gin Arg Pro Thr Asp Cys
- 50 His Leu Cys Gly Asp Ala Val Pro Arg Arg 165
- (2) INFORMATION FOR SEQ ID NO: 5: 55

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		(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 5:													
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20	Asp	Суѕ	Arg	Pro	Arg	Phe	Thr	Thr	Thr	Pro	Pro	Thr	Thr	Thr	Arg	Pro
20			115					120					125			
25	Pro	Arg 130	Arg	Arg	Arg											
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		(i)	SEG	20ENG	JE CI	HARA(	JIEK.	STIC	JS:							
30		( + )	(A) (B) (D)	) LI	ENGTI YPE:	H:	JIEK.	15110	i	148 a amine	o ac	o aci id	ids			
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35		(ii) (xi)	(A) (B) (D) MO)	) Li ) Ti ) To LECUI	ENGTE YPE: OPOLO LE T' CE DI	H: DGY: YPE: ESCRI	IPTI(	ON: S	SEQ :	amino linea Prote	o ac: ar ein O:	id 7:		Leu	Ile 15	Cys
	Met 1	(ii) (xi) Lys	(A) (B) (D) MOI SE(	) LH ) TO LECUI	ENGTH YPE: OPOLO LE T' CE DI Ala 5	H: DGY: YPE: ESCRI	IPTI( Leu	ON: S Gln	SEQ :	amino line Prote ID No Val 10	o ac ar ein O: Val	id 7: Ala	Leu	Leu Ser 30	15	
35	Met 1 Met	(ii) (xi) Lys Tyr	(A) (B) (D) MOI SE( Leu Asn	LECUI DUENG Thr	ENGT: YPE: OPCLO LE T' CE DI Ala 5	H: DGY: YPE: ESCRI Thr	IPTI( Leu Cys	ON: S Gln Val	SEQ : Val Ser 25	amino line Prote ID No Val 10 Gln	o actoring to a control of the contr	id 7: Ala Asn	Leu Asp	Ser	15 Pro	Pro
35 40	Met 1 Met Ser	(ii) (xi) Lys Tyr	(A) (B) (D) MOI SE( Leu Asn Asn 35	LECUI DUENG Thr Leu 20	ENGTHYPE: OPOLO LE T' CE DI Ala 5 Pro	H: DGY: YPE: ESCR: Thr Glu Met	IPTI( Leu Cys Arg	ON: S Gln Val Thr	SEQ : Val Ser 25	amino line. Prote ID No Val 10 Gln Asp	o ac: ar ein O: Val Ser	7: Ala Asn Ser	Leu Asp Gly 45	Ser 30	15 Pro Lys	Pro
35 40	Met 1 Met Ser	(ii) (xi) Lys Tyr Thr Asp	(A) (B) (D) MOD SEC Leu Asn Asn 35	LECUI DUENG Thr Leu 20 Asp	ENGTHYPE: OPOLO LE T' CE DI Ala 5 Pro Trp	H: DGY: YPE: ESCRI Thr Glu Met	Leu Cys Arg	ON: S Gln Val Thr 40 Gly	SEQ : Val Ser 25 Leu	amino line. Prote ID No Val 10 Gln Asp	o ac: ar ein O: Val Ser Lys	7: Ala Asn Ser Pro	Leu Asp Gly 45	Ser 30 Cys	Pro Lys	Pro Pro Asn
<b>35</b> <b>40</b> <b>45</b>	Met 1 Met Ser Arg Leu 65	(ii) (xi) Lys Tyr Thr Asp 50 Gln	(A) (B) (D) MOD SEC Leu Asn Asn 35 Thr	LECUI DUENG Thr Leu 20 Asp	ENGTHYPE: OPOLO LE T' CE DI Ala 5 Pro Trp Val	H: DGY: YPE: ESCRI Thr Glu Met Tyr Arg 70	Leu Cys Arg Leu 55 Cys	ON: S Gln Val Thr 40 Gly Val	SEQ : Val Ser 25 Leu Glu	amino line. Prote ID No Val 10 Gln Asp Val Val	ein O: Val Ser Lys Tyr Lys 75	7: Ala Asn Ser Pro 60 Arg	Leu Asp Gly 45 Glu Cys	Ser 30 Cys	Pro Lys Thr	Pro Pro Asn Cys

		130					135					140				
_	Glu 145	Pro	Arg	Arg												
5	(2)	INFO	RMAT	CION	FOR	SEQ	1D N	0:	8:							
		(i)	SEC	OUENC	CE CH	iarac	TERI	STIC	CS:							
10			(A) (B) (D)	TY	ENGTH (PE: OPOLO				ě	160 a amino linea	aci		ids			
15		(ii)	MOI	LECUI	LE TY	PE:			,	Prote	ein					
13		(xi)	SEÇ	QUENC	CE DI	ESCRI	PTIC	: : : : : : : : : : : : : : : : : : :	SEQ :	ID NO	): {	3:				
20	Pro 1	Ser	His	Gln	Lys 5	Lys	Val	Val	Pro	Trp 10	Ile	Asp	Val	Tyr	Thr 15	Arg
2.0	Ala	Thr	Cys	Gl n 20	Pro	Arg	Glu	Val	Val 25	Val	Pro	Leu	Sor	Mot 30	Glu	Leu
25	Met	Gly	Asn 35	Val	Val	Lys	Gln	Leu 40	Val	Pro	Ser	Суѕ	Val 45	Thr	Val	Gln
	Arg	Cys 50	Gly	Gly	Cys	Cys	Pro 55	Asp	Asp	Gly	Leu	Glu 60	Cys	Val	Pro	Thr
30																
	Gly 65	Gln	His	Gln	Val	Arg 70	Met	Gln	Ile	Leu	Met 75	Ile	Gln	Tyr	Pro	Ser 80
35	Ser	Gln	Leu	Gly	Glu 85	Met	Ser	Leu	Glu	Glu 90	His	Ser	Gln	Cys	Glu 95	Cys
40	Arg	Pro	Lys	Lys 100	Lys	Glu	Ser	Ala	Val 105	Lys	Pro	Asp	Ser	Pro 110	Arg	Ile
10	Leu	Cys	Pro 115	Pro	Cys	Thr	Gln	Arg 120	Arg	Gln	Arg	Pro	Asp 125	Pro	Arg	Thr
45	Cys	Arg 130	Cys	Arg	Cys	Arg	Arg 135		Arg	Phe	Leu	His 140		Gln	Gly	Arg
	Gly 145		Glu	Leu	Asn	Pro 150	Asp	Thr	Cys	Arg	Cys 155	Arą	Lys	Pro	Arq	Lys 160
50																
	(2)	INF	ORMA'	TION	FOR	SEQ	ID	:СИ	9:							
55		(±)	SE	QUEN	CE C	HARA:	CTER	ISTI	cs:							
55			4,5	, i	मश्या	н:				10,5	; .					

	Lys 1	Val	Val	Pro	Trp 5	Ile	Asp	Val	Tyr	Thr 10	Arg	Ala	Thr	Cys	Gln 15	Pro
5	Arg	Glu	Val	Val 20	Val	Pro	Leu	Ser	Met 25	Glu	Leu	Met	Gly	Asn 30	Val	Val
10	Lys	Gln	Leu 35	Val	Pro	Ser	Cys	Val 40	Thr	Val	Gln	Arg	Cys 45	Gly	Gly	Cys
10	Cys	Pro 50	Asp	Asp	Gly	Leu	Glu 55	Cys	Val	Pro	Thr	Gly 60	Gln	His	Gln	Val
15	Arg 65	Met	Gln	Ile	Leu	Met 70	Ile	Gln	Tyr	Pro	Ser 75	Ser	Gln	Leu	Gly	Glu 80
	Met	Ser	Leu	Glu	Glu 85	His	Ser	Gln	Cys	Glu 90	Cys	Arg	Pro	Lys	Lys 95	Lys
20	Glu	Ser	Ala	Val 100	Lys	Pro	Asp	Ser	Pro 105	Arg	Ile	Leu	Суѕ	Pro 110	Pro	Cys
25	Thr	Gln	Arg 115	Arg	Gln	Arg	Pro	Asp 120	Pro	Arg	Thr	Cys	Arg 125	Cys	Arg	Cys
_ •	Arg	Arg 130	Arg	Arg	Phe	Leu	His 135	Cys	Gln	Gly	Arg	Gly 140	Leu	Glu	Leu	Asn
30	Pro 145	Asp	Thr	Суѕ	Arg	Cys 150	Arg	Lys	Pro	Arg	Lys 155					
	(2)	INF	ORMA!	TION	FOR	SEQ	ID 1	10:	10:							
35		(i)	SE	QUENC	CE C	HARA	CTER	ISTI	CS:							
			(A) (B) (D)	Т.	ENGTI YPE: OPOLO				č	152 a amino linea	o ac.		ids			
40		(ii)	MOI	LECUI	LE T	YPE:			1	Prote	ein					
		(xi)	SE	QUEN	CE DI	ESCR:	IPTI	ON:	SEQ :	ID N	0:	10:				
45	Pro 1	Trp	Ile	Asp	Val 5	Tyr	Thr	Arg	Ala	Thr 10	Cys	Gln	Pro	Arg	Glu 15	Va:
	Val	Val	Pro	Leu 20	Ser	Met	Giu	Leu	Met 25	Gly	Arn	Val	Vāl	Lys 30	G1:.	Leu
50	Val	Pro	Ser 35	Cys	Val	Thr	Val	Gln 40	Arg	Cys	Gly	G1 y	Cys 45	Cys	Pro	Asp
<b>5</b> 5	Asp	Gly 50	Leu	Jlu	Cys	Val	Pro 55	Thr	Gly	Gln	His	Gln 60	Val	Arg	Met	Gln
	Ţ þ o	Len	Mr. t	π1 <sub>7</sub> ,	713 ·	············	Tive :		· ,-	A .	-			• • •		÷

				100					105					110		
5	Arg	Gln	Arg 115	Pro	Asp	Pro	Arg	Thr 120	Суз	Arg	Cys	Arg	Cys 125	Arg	Arg	Arg
3	Arg	Phe 130	Leu	His	Cys	Gln	Gly 135	Arg	Gly	Leu	Glu	Leu 140	Asn	Pro	Asp	Thr
10	Cys 145	Arg	Cys	Arg	Lys	Pro 150	Arg	Lys								
<b>1</b> 5	(2)	INFO	ORMAT	гіои	FOR	SEQ	ID 1	10:	11:							
		(i)	SEÇ	QUENC	CE CH	HARA(	CTERI	STI	CS:							
20			(A) (B) (D)	T	ENGTH (PE: OPOLO					150 a amino linea	ac		ids			
		(ii)	MOI	LECUI	LE TY	PE:				Prote	ein					
0.5		(xi)	SEÇ	QUENC	CE DE	ESCR:	IPTI(	)N: S	SEQ	ID NO	): :	11:				
25	Ile 1	Asp	Val	Tyr	Thr 5	Arg	Ala	Thr	Cys	Gln 10	Pro	Arg	Glu	Val	Val 15	Val
30	Pro	Leu	Ser	Met 20	Glu	Leu	Met	Gly	Asn 25	Val	Val	Lys	Gln	Leu 30	Val	Pro
	Ser	Cys	Val 35	Thr	Val	Gln	Arg	Cys 40	Gly	Gly	Cys	Cys	Pro 45	Asp	Asp	Gly
35	Leu	Glu 50	Суѕ	Val	Pro	Thr	Gly 55	Gln	His	Gln	Val	Arg 60	Met	Gln	Ile	Leu
40	Met 65	Ile	Gln	Tyr	Pro	Ser 70	Ser	Gln	Leu	Gly	Glu 75	Met	Ser	Leu	Glu	Glu 80
40	His	Ser	Gln	Cys	Glu 85	Cys	Arg	Pro	Lys	Lys 90	Lys	Glu	Ser	Ala	Val 95	Lys
45	Pro	Asp	Ser	Pro 100	Arg	Ile	Leu	Cys	Pro 105	Pro	Cys	Thr	Gln	Arg 110	Arg	Gln
	Arg	Pro	Asp 115	Pro	Arg	Thr	Cys	Arg 120	Cys	Arg	Cys	Arg	Arg 125	Arg	Ārg	Phe
50	Leu	His 130	Cys	Gln	Gly	Arg	Gly 135	Leu	Glu	Leu	Asn	Pro 140	Asp	Thr	Cys	Arg
55	Cys 145	Arg	Lys	Pro	Arg	Lys 150										
55	100	INFO	ORMA!	rrori	コヘロ	250	÷ r		· · .							

		(ii)	MOI	LECUI	E TY	PE:			ì	°rot€	ein					
Е		(xi)	SEÇ	QUENC	CE DE	ESCRI	PTIC	ON: S	SEQ I	D NO	): J	.2:				
5	Tyr 1	Thr	Arg	Ala	Thr 5	Суѕ	Gln	Pro	Arg	Glu 10	Val	Val	Val	Pro	Leu 15	Ser
10	Met	Glu	Leu	Met 20	Gly	Asn	Val	Val	Lys 25	Gln	Leu	Val	Pro	Ser 30	Cys	Val
	Thr	Val	Gln 35	Arg	Суѕ	Gly	Gly	Cys 40	Cys	Pro	Asp	Asp	Gly 45	Leu	Glu	Cys
15	Val	Pro 50	Thr	Gly	Gln	His	Gln 55	Val	Arg	Met	Gln	Ile 60	Leu	Met	Ile	Gln
20	Tyr 65	Pro	Ser	Ser	Gln	Leu 70	Gly	Glu	Met	Ser	Leu 75	Glu	Glu	His	Ser	Gln 80
	Cys	Glü	Cys	Arg	Pro 85	Lys	Lys	Lys	Glu	Ser 90	Ala	Val	Lys	Orq	Asp 95	Ser
25	Pro	Arg	Ile	Leu 100	Cys	Pro	Pro	Cys	Thr 105	Gln	Arg	Arg	Gln	Arg 110	Pro	Asp
	Pro	Arg	Thr 115	Cys	Arg	Cys	Arg	Cys 120	Arg	Arg	Arg	Arg	Phe 125	Leu	His	Cys
30	Gln	Gly 130	Arg	Gly	Leu	Glu	Leu 135	Asn	Pro	Asp	Thr	Cys 140	Arg	Cys	Arg	Lys
35	Pro 1 <b>4</b> 5	Arg	Lys													
	(2)	INF	ORMA	TION	FOR	SEQ	ID I	: 00	13:							
40		(i)	SE	QUEN	CE C	HARA(	CTER:	ISTI	CS:							
			(A (B (D	) T	ENGTI YPE: OPOLO				•		o ac	o ac: id	ids			
45		(i1)	MO:	LECU:	LE T	YPE:				Frot	ein					
		(xi)	SE	QUEN(	GE D	ESCR	IPTI	: NC	SEQ	ID N	0:	13:				
50	Arg 1	Ala	Thr	Cys	Gln 5	Pro	Arg	Glu	Val	Vai 10	Val	Pro	Leu	Ser	Met 15	Glu
	Ľeu	Met	Gly	Asn 20	Val	Val	Lys	Gln	Leu 25	Val	Pro	Ser	Cys	Val 30	Thr	Val
55	Gln	Arg	Cys	Gly	Gly	Cys	Cys	Pro	Asp	Asp	Gly	Leu	Glu	Cys	Va!	Pro

ing the second of the second o

	Cys	Arg	Pro	Lys	Lys 85	Lys	Glu	Ser	Ala	Val 90	Lys	Pro	Asp	Ser	Pro 95	Arg
5	Ile	Leu	Cys	Pro 100	Pro	Cys	Thr	Gln	Arg 105	Arg	Gln	Arg	Pro	Asp 110	Pro	Arg
10	Thr	Суз	Arg 115	Суѕ	Arg	Cys	Arg	Arg 120	Arg	Arg	Phe	Leu	His 125	Cys	Gln	Gly
10	Arg	Gly 130	Leu	Glu	Leu	Asn	Pro 135	Asp	Thr	Cys	Arg	Cys 140	Arg	Lys	Pro	Arg
15	Lys 145 (2)	INF	ORMAT	TION	FOR	SEQ	ID 1	10:	14:							
		(i)	SEÇ	QUENC	CE CI	HARAC	CTER	STIC	cs:							
20			(A) (B) (D)	т (	ENGTI YPE: DPOLO				3	178 a amino linea	o aci		ids			
25		(ii)	MOI	LECUI	LE T	YPE:			1	Prote	ein					
<b>2</b> J		(xi)	SEÇ	QUEN	CE DI	ESCR:	PTIC	: : NC	SEQ :	ID NO	): :	L4:				
30	Pro 1	Gly	His	Gln	Arg 5	Lys	Val	Val	Ser	Trp 10	Ile	Asp	Val	Tyr	Thr 15	Arg
50	Ala	Thr	Cys	Gln 20	Pro	Arg	Glu	Val	Val 25	Val	Pro	Leu	Thr	Val 30	Glu	Leu
35	Met	Gly	Thr 35	Val	Ala	Lys	Gln	Leu 40	Val	Pro	Ser	Cys	Val 45	Thr	Val	Gln
	Arg	Cys 50	Gly	Gly	Cys	Cys	Pro 55	Asp	qzA	Gly	Leu	Glu 60	Cys	Val	Pro	Thr
40	Gly 65	Gln	His	Gln	Val	Arg 70	Met	Gln	Ile	Leu	Met 75	Ile	Arg	Tyr	Pro	Ser 80
45	Ser	Gln	Leu	Gly	Glu 85	Met	Ser	Leu	Glu	Glu 90	His	Ser	Gln	Cys	Glu 95	Cys
10	Arg	Pro	Lys	Lys 100	Asp	Ser	Ala	Val	195 105	Pro	Asp	Arg	Alā	Ala 110	Thr	Pro
50	His	His	Arg 115	Fro	Glr.	Pro	Arg	Ser 120	Val	Pro	Gly	Trp	Asp 125	ser	Ala	Pro
	Gly	Ala 130		Ser	Pro	Ala	Asp 135		Thr	His	Pro	Thr 140	Pro	λla	Pro	GIA
55	Fro 145		Ala	His	Ala	Ala	Pro	Ser	Tnr	Thr	Ser	Ala	Leu	Thr	Pro	gly

	(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	IC:	15:							
		( <u>i</u> )	SEÇ	QUENC	CE CH	IARAC	CTERI	STIC	S:							
5			(A) (B) (D)	TY	NGTH PE: POLC					173 ā amino lineā	aci		.ds			
10		(ii)	MOI	LECUI	E TY	PE:				Prote	ein					
10		(xi)	SEÇ	QUENC	CE DE	ESCRI	PTIC	on: s	SEQ	ID NO	): 1	15:				
15	Lys 1	Val	Val	Ser	Trp 5	Ile	Asp	Val	Tyr	Thr 10	Arg	Ala	Thr	Cys	Gln 15	Pro
15	Arg	Glu	Val	Val 20	Val	Pro	Leu	Thr	Val 25	Glu	Leu	Met	Gly	Thr 30	Val	Ala
20	Lys	Gln	Leu <b>3</b> 5	Val	Pro	Ser	Суѕ	Val 40	Thr	Val	Gln	Arg	Cys 45	Gly	Gly	Cys
	Cys	Pro 50	Asp	Asp	Gly	Leu	Glu 55	Суз	Val	Pro	Thr	Gly 60	Gln	His	Gln	Val
25	Arg 65	Met	Gln	Ile	Leu	Met 70	Ile	Arg	Tyr	Pro	Ser 75	Ser	Gln	Leu	Gly	Glu 80
30	Met	Ser	Leu	Glu	Glu 85	His	Ser	Gln	Cys	Glu 90	Cys	Arg	Pro	Lys	Lys 95	Asp
	Ser	Ala	Val	Lys 100	Pro	Asp	Arg	Ala	Ala 105	Thr	Pro	His	His	Arg 110	Pro	Gln
35	Pro	Arg	Ser 115	Val	Pro	Gly	Trp	Asp 120	Ser	Ala	Pro	Gly	Ala 125	Pro	Ser	Pro
	Ala	Asp 130		Thr	His	Pro	Thr 135	Pro	Ala	Pro	Gly	Pro 140	Ser	Ala	His	Ala
40	Ala 145		Ser	Thr	Thr	Ser 150	Ala	Leu	Thr	Pro	Gly 155	Pro	Ala	Ala	Ala	Ala 160
45	Ala	Asp	Ala	Ala	Ala 165		Ser	Val	Ala	Lys 170	Gly	Gly	Ala			
	(2)	INF	ORMA'	TION	FOR	SEQ	ID :	NO:	16:							
		(1)	SE	QUEN	CE C	HARA	CTER	ISTI	CS:							
50			(B	) L: ) T:	YPE:					168 amin line	o ac		ids			
		(±i)	MO	LECU	LE T	YPE:				Prot	ein					
55		/ <sub>X</sub> :	10			5035	T.F.M.	· · · .		~~ ,,		ē				

	Ser	Cys	Val 35	Thr	Val	Gln	Arg	Cys 40	Gly	Gly	Cys	Cys	Pro 45	Asp	Asp	Gly
5	Leu	Glu 50	Cys	Val	Pro	Thr	Gly 55	Gln	His	Gln	Val	Arg 60	Met	Gln	Ile	Leu
1.0	Met 65	Ile	Arg	Tyr	Pro	Ser 70	Ser	Gln	Leu	Gly	Glu 75	Met	Ser	Leu	Glu	Glu 80
10	His	Ser	Gln	Cys	Glu 85	Cys	Arg	Pro	Lys	Lys 90	Asp	Ser	Ala	Val	Lys 95	Pro
15	Asp	Arg	Ala	Ala 100	Thr	Pro	His	His	Arg 105	Pro	Gln	Pro	Arg	Ser 110	Val	Pro
	Gly	Trp	Asp 115	Ser	Ala	Pro	Gly	Ala 120	Pro	Ser	Pro	Ala	Asp 125	Ile	Thr	His
20	Pro	Thr 130	Pro	Ala	Pro	Gly	Pro 135	Ser	Ala	His	Ala	Ala 140	Pro	Ser	Thr	Thr
25	Ser 145	Ala	Leu	Thr	Pro	Gly 150	Pro	Ala	Ala	Ala	Ala 155	Ala	Asp	Ala	Ala	Ala 160
23	Ser	Ser	Val	Ala	Lys 165	Gly	Gly	Ala								
30	(2)	INF	ORMA!	NOIT	FOR	SEQ	ID 1	10:	17:							
		(i)	SEQ	QUENC	CE C	IARA(	CTER	STIC	CS:							
35			(A) (B) (D)	) T	ENGTI YPE: OPOLO				á		o ac	o ac: id	ds			
		(ii)	MOI	LECUI	LE TY	YPE:			I	Prote	ein					
4.0		(xi)	SEQ	QUEN	CE DE	ESCR	IPTIC	ON: 8	SEQ I	ID NO	): :	17:				
40	Arg 1	Ala	Thr	Cys	Gln 5	Pro	Arg	Glu	Val	Val	Val	Pro	Leu	Thr	Val 15	Glu
45	Leu	Met	Gly	Thr 20			Lys		Leu 25	Val	Pro	Ser	Cys	Val 30	Thr	Val
	Gln	Arg	Jys 35	Gly	Gly	Cys	Cys	Pro 40	Asp	Asp	Gly	Leu	Glu 45	Суз	Val	Pro
50	Thr	Gly 50	Gln	His	Gln	Val	Λrg 55	Met	Gln	Ile	Leu	Met 60	Ile	Λrg	Tyr	Pro
55	Ser 65	Ser	Gln	Leu	Gly	Glu 70	Met	Ser	Leu	Glu	Glu 75	His	Ser	Gln	Cys	Glu 80
J J	Cvs	Ara	Pro	Lus	T ++1+	A	c	7,1 .	• • • •	r	5	: .	٠			

			115					120					125			
5	Gly	Pro 130	Ser	Ala	His	Ala	Ala 135	Pro	Ser	Thr	Thr	Ser 140	Ala	Leu	Thr	Pro
J	Gly 145	Pro	Ala	Ala	Ala	Ala 150	Ala	Asp	Ala	Ala	Ala 155	Ser	Ser	Val	Ala	Lys 160
10	Gly	Gly	Ala													
- 0	(2)	INF	ORMA'	NOIT	FOR	SEQ	ID	10:	18:							
		(i)	SEÇ	QUENC	CE C	HARA	CTER:	ISTIC	CS:							
15			(A) (B) (D)	T	ENGTI YPE: OPOLO				ő	194 a amino linea	ac:		ids			
20		(ii)	MOI	LECUI	LE T	YPE:			I	Prote	ein					
20		(xi)	SEÇ	QUENC	TE DI	ESCRI	IPTI	: MC	SEQ :	ID NO	); ]	18:				
25	Pro 1	Gly	His	Gln	Arg 5	Lys	Val	Val	Ser	Trp 10	Ile	Asp	Val	Tyr	Thr 15	Arg
	Ala	Thr	Суѕ	Gln 20	Pro	Arg	Glu	Val	Val 25	Val	Pro	Leu	Thr	Val 30	Glu	Leu
30	Met	Gly	Thr 35	Val	Ala	Lys	Gln	Leu 40	Val	Pro	Ser	Cys	Val 45	Thr	Val	Gln
	Arg	Cys 50	Gly	Gly	Суѕ	Cys	Pro 55	Asp	Asp	Gly	Leu	Glu 60	Cys	Val	Pro	Thr
35	Gly 65	Gln	His	Gln	Val	Arg 70	Met	Gln	Ile	Leu	Met 75	Ile	Arg	Tyr	Pro	Ser 80
40	Ser	Gln	Leu	Gly	Glu 85	Met	Ser	Leu	Glu	Glu 90	His	Ser	Gln	Суѕ	Glu 95	Cys
. •	Arg	Pro	Lys	Lys 100	րչ	Asp	Ser	Ala	Val 105	Lys	Gln	Лsp	Arg	Ala 110	Ala	Thr
45	Pro	His	His 115	Arg	Pro	Gln	Pro	Arg 120	Ser	Val	Pro	Gly	Trp 125	Asp	Ser	Ala
	Pro	Gly 130	Ala	Pro	Ser	Pro	Λla 135	Λsp	He	Thr	Gln	Ser 140	His	Ser	Set	Pro
50	Arg 145	Pro	Leu	Cys	Pro	Arg 150	Cys	Thr	Gln	His	His 155	Gln	Cys	Pro	Asp	Pro 160
55					165					170			Leu		175	
	Glv	Arg	ai.	Ton	ينت	÷ , , ,	7	77 5-			~	٠, .		٠,	•	•

		(i)	SEÇ	QUENC	CE CI	iara(	CTER	STIC	CS:							
5			(A) (B) (D)	TY	ENGTH (PE: )POL(					189 a amino linea	ac:		ds			
		(ii)	MOI	LECUI	LE TY	PE:				Prote	ein					
10		(xi)	SEÇ	QUENC	CE DE	ESCR	PTIC	ON: 5	SEQ	ID NO	): [	19:				
10	Lys 1	Val	Val	Ser	Trp 5	Ile	Asp	Val	Tyr	Thr 10	Arg	Ala	Thr	Cys	Gln 15	Pro
15	Arg	Glu	Val	Val 20	Val	Pro	Leu	Thr	Val 25	Glu	Leu	Met	Gly	Thr 30	Val	Ala
	Lys	Gln	Leu 35	Val	Pro	Ser	Cys	Val 40	Thr	Val	Gln	Arg	Cys 45	Gly	Gly	Cys
20	Суѕ	Pro 50	Asp	Asp	Gly	Leu	Glu 55	Cys	Val	Pro	Thr	Gly 60	Gln	His	Gln	Val
25	Arg 65	Met	Gln	Ile	Leu	Met 70	Ile	Arg	Tyr	Pro	Ser 75	Ser	Gln	Leu	Gly	Glu 80
20	Met	Ser	Leu	Glu	Glu 85	His	Ser	Gln	Cys	Glu 90	Cys	Arg	Pro	Lys	Lys 95	Lys
30	Asp	Ser	Ala	Val 100	Lys	Gln	Asp	Arg	Ala 105	Ala	Thr	Pro	His	His 110	Arg	Pro
	Gln	Pro	Arg 115	Ser	Val	Pro	Gly	Trp 120	Asp	Ser	Ala	Pro	Gly 125	Ala	Pro	Ser
35	Pro	Ala 130	Asp	Ile	Thr	Gln	Ser 135	His	Ser	Ser	Pro	Arg 140	Pro	Leu	Cys	Pro
40	Arg 145	Cys	Thr	Gln	His	His 150	Gln	Cys	Pro	Asp	Pro 155	Arg	Thr	Cys	Arg	Cys 160
.0	Arg	Cys	Arg	Arg	Arg 165	Ser	Phe	Leu	Arg	Cys 170	Gln	Gly	Arg	Gly	Leu <b>1</b> 75	Glu
45	Leu	Asn	Pro	Asp 180	Thr	Cys	Arg	Cys	Arg 185	Lys	Leu	Arg	Arg			
	(2)	INFO	ORMA!	иол	FOR	SEQ	ID I	: ⊊и	20:							
50		(1)	SEÇ	QUEN	CE C	IARA	CTER.	ISTI	CS:							
30			(A) (B) (D)		ENGTI YPE: OPOLO					184 a amine line	o ac		ids			
55		(iı)	MOI	LECU:	LE T	YPE:				Prote	ein					

				20					25					30		
r	Ser	Cys	Val 35	Thr	Val	Gln	Arg	Cys 40	Gly	Gly	Cys	Cys	Pro	Asp	Asp	Gly
5	Leu	Glu 50	Cys	Val	Pro	Thr	Gly 55	Gln	His	Gln	Val	Arg 60	Met	Gln	Ile	Leu
10	Met 65	Ile	Arg	Tyr	Pro	Ser 70	Ser	Gln	Leu	Gly	Glu 75	Met	Ser	Leu	Glu	Glu 80
	His	Ser	Gln	Cys	Glu 85	Cys	Arg	Pro	Lys	Lys 90	Lys	Asp	Ser	Ala	Val 95	Lys
15	Gln	Asp	Arg	Ala 100	Ala	Thr	Pro	His	His 105	Arg	Pro	Gln	Pro	Arg 110	Ser	Val
20	Pro	Gly	Trp 115	Asp	Ser	Ala	Pro	Gly 120	Ala	Pro	Ser	Pro	Ala 125	Asp	Ile	Thr
20	Gln	Ser 130	His	Ser	Ser	Pro	Arg 135	Pro	Leu	Суз	Pro	Ary 140	Суз	Thr	Gln	His
25	His 145	Gln	Суз	Pro	Asp	Pro 150	Arg	Thr	Cys	Arg	Cys 155	Arg	Cys	Arg	Arg	Arg 160
	Ser	Phe	Leu	Arg	Cys 165	Gln	Gly	Arg	Gly	Leu 170	Glu	Leu	Asn	Pro	Asp 175	Thr
30	Cys	Arg	Cys	Arg 180	Lys	Leu	Arg	Arg								
	(2)	INFO	ORMA'	rion	FOR	SEQ	ID	NO:	21:							
35		(i)	SE	QUEN	CE C	HARA	CTER:	ISTI	CS:							
			(A (B (D	) T	ENGT: YPE: OPOL				ě	179 a amine	o ac		ids			
40		(ii)	MO.	LECU:	LE T	YPE:				Prot	ein					
		(xi)	SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:	21:				
45	Arg 1	Al a	Thr	Cys	Gln 5	Pro	Arg	Glu	Vāl	Va! 10	Val	Pro	Lea	Thr	Val 15	Glu
5.0	Leu	Met	Gly	Thr 20	Val	Ala	Ъуs	Gln	Leu 25	Val	Pro	Ser	Cys	Val 30	Thr	Val
50	Gln	Arg	Cys 35	Gly	Gly	Cys	Суѕ	Pro 40	Asp	Asp	Gly	Leu	Gl u 45	Cys	Val	Pro
55	Thr	Gly 50	Gln	His	Gln	Val	Arg 55	Met	Gln	Tie	Leu	Met 60	Ile	Arg	Tyr	Pro

	Thr	Pro	His	His 100	Arg	Pro	Gln	Pro	Arg 105	Ser	Val	Fro	Gly	110		Ser
5	Ala	Pro	Gly 115	Ala	Pro	Ser	Pro	Ala 120	Asp	Ile	Thr	Gln	Ser 125	His	Ser	Ser
	Pro	Arg 130	Pro	Leu	Cys	Pro	Arg 135	Суѕ	Thr	Gln	His	Fis 140	Gln	Cys	Pro	Asp
10	Pro 145	Arg	Thr	Cys	Arg	Cys 150	Arg	Cys	Arg	Arg	Arg 155	Ser	Phe	Leu	Arg	Cys 160
15	Gln	Gly	Arg	Gly	Leu 165	Glu	Leu	Asn	Pro	Asp 170	Thr	Cys	Arg	Cys	Arg 175	Lys
	Leu	Arg	Arg													
	(2)	INF	ORMA'	NOIT	FOR	SEQ	I DI	10:	22:							
20		(i)	SEC	QUENC	CE CI	HARAC	CTER	ISTIC	CS:							
25			(A) (B) (D)	) T	ENGTI YPE: DPOLO				č	307 a amino linea	ac:		ids			
23		(ii)	MO]	LECUI	LE T	YPE:			I	Prote	ein					
		(xi)	SE	QUEN	CE DI	ESCR	IPTIC	ON: S	SEQ I	ID 110	); ;	22:				
30	His 1	Tyr	Asn	Thr	Glu 5	Ile	Leu	Lys	Ser	Ile 10	Asp	Asn	Glu	Trp	Arg 15	Lys
35	Thr	Gln	Cys	Met 20	Pro	Arg	Glu	Val	Cys 25	Ile	Asp	Val	Gly	Lys 30	Glu	Phe
	Gly	Val	Ala 35	Thr	Asn	Thr	Phe	Phe 40	Lys	Pro	Pro	Cys	Val 45	Ser	Val	Tyr
40	Arg	Cys 50	Gly	Gly	Cys	Cys	Asn 55	Ser	Glu	Gly	Leu	Gln 60	Суѕ	Met	Asn	Thr
	Ser 65	Thr	Ser	Tyr	Leu	Ser 70	Lys	Thr	Leu	Phe	Glu 75	Ile	Thr	Val	Pro	Leu 80
45	Ser	Gln	Gly	Pro	Lys 85	Pro	Val	Thr	Ile	Ser 90	Phe	Ala	Asn	His	Thr 95	Ser
50	Cys	Arg	Суѕ	Met 100	Ser	Lys	Leu	Asp	Val 105		Arg	Gln	Val	His 110	Ser	Ile
	Ile	Arg	Arg 115	Ser	Leu	Pro	Ala	Thr 120		Pro	Gln	Cys	Gln 125		Ala	Asn
55	Lys	Thr 130		Pro	Thr	Asn	Туг 135		Trp	Asn	Λcn	His 140		Çys	Arg	Cys

	Glu	Thr	Cys	Gln 180	Cys	Val	Cys	Arg	Ala 185	Gly	Leu	Arg	Pro	Ala 190	Ser	Cys
5	Gly	Pro	His 195	Lys	Glu	Leu	Asp	Arg 200	Asn	Ser	Cys	Gln	Cys 205	Val	Cys	Lys
	Asn	Lys 210	Leu	Phe	Pro	Ser	Gln 215	Cys	G]y	Ala	Asn	Arg 220	Glu	Phe	Asp	Glu
10	Asn 225	Thr	Cys	Gln	Cys	Val 230	Cys	Lys	Arg	Thr	Cys 235	Pro	Arg	Asn	Gln	Pro 2 <b>4</b> 0
15	Leu	Asn	Pro	Gly	Lys 245	Cys	Ala	Cys	Glu	Cys 250	Thr	Glu	Ser	Pro	Gln 255	Lys
	Cys	Leu	Leu	Lys 260	Gly	Lys	Lys	Phe	His 265	His	Gln	Thr	Cys	Ser 270	Cys	Tyr
20	Arg	Arg	Pro 275	Cys	Thr	Asn	Arg	Gln 280	Lys	Ala	Cys	Glu	Pro 285	Gly	Phe	Ser
	Tyr	Ser 290	Glu	Glu	Val	Cys	Arg 295	Cys	Val	Pro	Ser	Tyr 300	Trp	Lys	Arg	Pro
25	Gln 305	Met	Ser													
	(2)	INF	ORMA?	rion	FOR	SEQ	ID i	:07	23:							
30		(i)	SE	QUEN	CE C	HARA(	CTER:	ISTIC	CS:							
<b>3</b> E			(A) (B) (D)	) Т	ENGT: YPE: OPOLO				ä	302 a amino linea	ac:	o ac. id	ids			
35		(ii)	MOI	LECU	LE T	YPE:			1	Prote	ein					
		(xi)	SE	QUEN	CE DI	ESCR:	IPTI	): :	SEQ :	ID N	): :	23:				
40	Ile 1	Leu	Lys	Ser	Ile 5	Asp	Asn	Glu	Trp	Arg 10	Lys	Thr	Gln	Cys	Met 15	Pro
<b>4</b> 5	Arg	Glu	Val	Cys 20	Ile	Asp	Val	Gly	Lys 25	Glu	Phe	Gly	Val	Ala 30	Thr	Asn
<b>4</b> 5	Thr	Phe	Phe 35	Lys	Pro	Pro	Cys	Val 40	Ser	Val	Tyr	Arg	Cys 45	Gly	Gly	Cys
50	ੇys	Asn 50	Ser	Glu	Gly	Leu	Gln 55	Cys	Met	Asn	Thr	Ser 60	Thr	šer	Tyr	Leu
	Ser 65	Lys	Thr	Leu	Phe	Glu 70	Ile	Thr	Val	Pro	Leu 75	Ser	Gln	31 y	Pro	Lys 80
55	Pro	Val	Thr	Ile	Ser	Phe	Ala	Asn	Eis	Thr	Ser	Cys	Arq	Cys	Met	Ser

<sup>115</sup> II We will have all him lyr that tyr in 116 115

	Asn	Tyr 130	Met	Trp	Asn	Asn	His 135	Ile	Cys	Arg	Cys	Leu 140	Ala	Gln	Glu	Asp
5	Phe 145	Met	Phe	Ser	Ser	Asp 150	Ala	Gly	Asp	Asp	Ser 155	Thr	Asp	Gly	Phe	His 160
10	Asp	Ile	Cys	Gly	Pro 165	Asn	Lys	Glu	Leu	Asp 170	Glu	Glu	Thr	Cys	Gln 175	Cys
<b>1</b> F	Val	Cys	Arg	Ala 180	Gly	Leu	Arg	Pro	Ala 185	Ser	Cys	Gly	Pro	His 190	Lys	Glu
15	Leu	Asp	Arg 195	Asn	Ser	Cys	Gln	Cys 200	Val	Cys	Lys	Asn	Lys 205	Leu	Phe	Pro
20	Ser	Gln 210	Суѕ	Gly	Ala	Asn	Arg 215	Glu	Phe	Asp	Glu	Asn 220	Thr	Cys	Gln	Cys
	Val 225	Cys	Lys	Arg	Thr	Cys 230	Pro	Arg	Asn	Gln	Pro 235	Leu	Asn	Pro	Gly	Lys 240
25	Cys	Ala	Cys	Glu	Cys 245	Thr	Glu	Ser	Pro	Gln 250	Lys	Cys	Leu	Leu	Lys 255	Gly
30	Lys	Lys	Phe	His 260	His	Gln	Thr	Cys	Ser 265	Cys	Tyr	Arg	Arg	Pro 270	Cys	Thr
50	Asn	Arg	Gln 275	Lys	Ala	Cys	Glu	Pro 280	Gly	Pł.e	Ser	Tyr	Ser 285	Glu	Glu	Val
35	Cys	Arg 290	Cys	Val	Pro	Ser	Tyr 295	Trp	Lys	Arg	Pro	Gln 300	Met	Ser		
	(2)	INF	ORMA:	rion	FOR	SEQ	ID	NO:	24:							
40		(i)	SEÇ	QUEN	CE C	HARA	CTER:	ISTI	CS:							
			(A) (B) (D)	T	ENGTI YPE: OPOLO				ć	297 a amino linea	o ac		ids			
45		(ii)	MO:	LECU:	LE T	YPE:				Prote	ein					
		(xi)	SE	QUEN	CE DI	ESCR	IFTI	DN: 1	SEC	ID N	.:: .	24:				
50	Asp 1	Asn	Glu	Trp	Arg 5	Lys	Thi	Gln	Cys	Met 10	Pro	Arg	Jiu	Val	Cys 15	lle
	Asp	Val	Gly	Lys 20	Glu	Phe	Cly	Val	Ala 25	Thr	Asn	Thr	Phe	Phe 30	Lys	Pro
55	Pro	Cys	Val 35	Ser	Val	Tyr	Arg	Cys	Gly	Gly	Cys	Cys	Asn	Ser	Glu	Gly

Fig. 1.6 In: Val. 1.6 (1944) + 1 (2011) 17 (2011) 19 (4) (1941) 18

	Phe	Ala	Asn	His	Thr 85	Ser	Cys	Arg	Cys	Met 90	Ser	Lys	Leu	Asp	Val 95	Tyr
5	Arg	Gln	Val	His 100	Ser	Ile	Ile	Arg	Arg 105	Ser	Leu	Pro	Ala	Thr 110	Leu	Pro
10	Gln	Cys	Gln 115	Ala	Ala	Asn	Lys	Thr 120	Cys	Pro	Thr	Asn	Tyr 125	Met	Trp	Asn
10	Asn	His 130	Ile	Суѕ	Arg	Cys	Leu 135	Ala	Gln	Glu	Asp	Phe 140	Met	Phe	Ser	Ser
15	Asp 145	Ala	Gly	Asp	Asp	Ser 150	Thr	Asp	Gly	Phe	His 155	Asp	Ile	Cys	Gly	Pro 160
	Asn	Lys	Glu	Leu	Asp 165	Glu	Glu	Thr	Cys	Gln 170	Cys	Val	Суѕ	Arg	Ala 175	Gly
20	Leu	Arg	Pro	Ala 180	Ser	Суѕ	Gly	Pro	His 185	Lys	Glu	Leu	Asp	Arg 190	Asn	Ser
25	Cys	Gln	Cys 195	Val	Cys	Lys	Asn	Lys 200	Leu	Phe	Pro	Ser	Gln 205	Cys	Gly	Ala
23	Asn	Arg 210	Glu	Phe	Asp	Glu	Asn 215	Thr	Cys	Gln	Cys	Val 220	Суѕ	Lys	Arg	Thr
30	Cys 225	Pro	Arg	Asn	Gln	Pro 230	Leu	Asn	Pro	Gly	Lys 235	Cys	Ala	Cys	Glu	Cys 240
	Thr	Glu	Ser	Pro	Gln 245	Lys	Cys	Leu	Leu	Lys 250	Gly	Lys	Lys	Phe	His 255	His
35	Gln	Thr	Cys	Ser 260	Cys	Tyr	Arg	Arg	Pro 265	Cys	Thr	Asn	Arg	Gln 270	Lys	Ala
<b>4</b> 0	Cys	Glu	Pro 275	Gly	Phe	Ser	Tyr	Ser 280	Glu	Glu	Val	Cys	Arg 285	Cys	Val	Pro
	Ser	Tyr 290	Trp	Lys	Arg	Pro	Gln 295	Met	Ser							
45	(2)		ORMA' SE(					NO: ISTIC	25: CS:							
50			(B)	T.		H: DGY:				292 amino	o ac.		ids			
		(ii)	MO	LECU:	LE T	YPE:				Prot	ein					
55		(xi)	SE	DUEN	CE DI	ESCR.	:PTI	ЭN: :	SEQ	ID N	·':	25:				
JŪ	Lys	Thr	Gln	عائان	No+	Pro	ār:	C * · ·	· ·	÷		٠.			• =	

			35					40					45			
5	Thr	Ser 50	Thr	Ser	Tyr	Leu	Ser 55	Lys	Thr	Leu	Phe	Glu 60	Ile	Thr	Val.	Pro
5	Leu 65	Ser	Gln	Gly	Pro	Lys 70	Pro	Val	Thr	Ile	Ser 75	Phe	Ala	Asn	His	Thr 80
10	Ser	Cys	Arg	Суѕ	Met 85	Ser	Lys	Leu	Asp	Val 90	Tyr	Arg	Gln	Val	His 95	Ser
	Ile	Ile	Arg	Arg 100	Ser	Leu	Pro	Ala	Thr 105	Leu	Pro	Gln	Суѕ	Gln 110	Ala	Ala
15	Asn	Lys	Thr 115	Cys	Pro	Thr	Asn	Tyr 120	Met	Trp	Asn	Asn	His 125	Ile	Cys	Arg
20	Cys	Leu 130	Ala	Gln	Glu	Asp	Phe 135	Met	Phe	Ser	Ser	Asp 140	Ala	Gly	Asp	Asp
	Ser 145	Thr	qaA	Gly	Phe	His 150	Asp	lle	Cys	GГУ	Pro 155	Asn	Lys	Glu	Leu	Asp 160
25	Glu	Glu	Thr	Cys	Gln 165	Cys	Val	Cys	Arg	Ala 170	Gly	Leu	Arg	Pro	Ala 175	Ser
	Cys	Gly	Pro	His 180	Lys	Glu	Leu	Asp	Arg 185	Asn	Ser	Cys	Gln	Cys 190	Val	Cys
30	Lys	Asn	Lys 195	Leu	Phe	Pro	Ser	Gln 200	Cys	Gly	Ala	Asn	Arg 205	Glu	Phe	Asp
35	Glu	Asn 210	Thr	Cys	Gln	Cys	Val 215	Cys	Lys	Arg	Thr	Cys 220	Pro	Arg	Asn	Gln
	Pro 225	Leu	Asn	Pro	Gly	Lys 230	Суѕ	Ala	Cys	Glu	Cys 235	Thr	Glu	Ser	Pro	Gln 240
40	Lys	Cys	Leu	Leu	Lys 245	Gly	Lys	Lys	Phe	His 250	His	Gln	Thr	Cys	Ser 255	Cys
	Tyr	Arg	Arg	Pro 260	Cys	Thr	Asn	Arg	Gln 265		Ala	Суѕ	Glu	Pro 270	Gly	Phe
45	Ser	Tyr	Ser 275	Glu	Glu	Val	Cys	Arg 280	Cys	Val	Pro	Ser	Tyr 285	Trp	Lys	Arç
50	Pro	Gln 290	Met	Ser												
	(2)	INF		TION												
		(i)	SE	QUEN	CE C	HARA	CTER	ISTI	CS:							
55			(A 'B		ENGT	Н:					amin		ids			

 $(1, \dots, 1, \dots, 1,$ 

	Leu 1	Asn	Ala	Asp	Ser 5	Asn	Thr	Lys	Gly	Trp 10	Ser	Glu	Val	Leu	Lys 15	G1 y
5	Ser	Glu	Cys	Lys 20	Pro	Arg	Pro	Ile	Val 25	Vāl	Pro	Val	Ser	Glu 30	Thr	His
	Pro	Glu	Leu 35	Thr	Ser	Gln	Arg	Phe 40	Asn	Pro	Pro	Суѕ	Val 45	Thr	Leu	Met
10	Arg	Cys 50	Gly	Gly	Cys	Cys	Asn 55	Asp	Glu	Ser	Leu	Glu 60	Cys	Val	Pro	Thr
1.5	Glu 65	Glu	Val	Asn	Val	Thr 70	Met	Glu	Leu	Leu	Gly 75	Ala	Ser	Gly	Ser	Gly 80
15	Ser	Asn	Gly	Met	Gln 85	Arg	Leu	Ser	Phe	Val 90	Glu	His	Lys	Lys	Cys 95	Asp
20	Cys	Arg	Pro	Arg 100	Phe	Thr	Thr	Thr	Pro 105	Pro	Thr	Thr	Thr	Arg 110	Pro	Pro
	Arg	Arg	Arg 115	Arg												
25																
	(2)	INF	ORMA!	rion	FOR	SEQ	ID 1	10:	27:							
		(i)	C E/													
30		(1)	SE(	20±N	CE C	HARAG	CTER.	ISTI	CS:							
30		(1)	(A (B (D	) L1	CE CI ENGT: YPE: OPOLO	н:	CTER.	ISTI(		111 amino	o ac	o ac id	ids			
30 35		(ii)	(A (B (D	) L1	ENGT: YPE: OPOL	H: OGY:	CTER.	ISTI		amin	o ac ar		ids			
			(A (B (D MO)	) LI ) T  ) TO	ENGT: YPE: OPOLO	H: OGY:				amine line Prot	o ac ar ein		ids			
		(ii) (xi)	(A) (B) (D) MO:	) LI ) T ) T ( LECU:	ENGT: YPE: OPOLO LE T	H: OGY: YPE: ESCR	IPTI	ON:	SEQ	amine line Prot	o ac ar ein O:	id 27:		Cys	Lys 15	Pro
35	Asn 1	(ii) (xi) Thr	(A (B (D MO: SEC	) LI ) T ) T ( LECU: QUEN ( Gly	ENGT: YPE: OPOL LE T CE D Trp	H: OGY: YPE: ESCR Ser	IPTI Glu	ON: : Val	SEQ Leu	emine line Prote ID N Lys 10	o ac ar ein O: Gly	id 27: Ser	Glu	Cys Leu 30	15	
35	Asn 1 Arg	(ii) (xi) Thr	(A (B (D MO) SEC Lys	) Li ) T' ) TO  LECU:  QUENC  Gly  Val 20	ENGT: YPE: OPOLO LE T CE D Trp 5	H: OGY: YPE: ESCR Ser Pro	IPTI Glu Val	ON: : Val Ser	SEQ Leu Glu 25	amino line Prote ID No Lys 10 Thr	o ac ar ein O: Gly His	id 27: Ser Pro	Glu	Leu	15 Thr	Ser
35 40 45	Asn 1 Arg	(ii) (xi) Thr Pro	(A) (B) (D) MO: SEC Lys Ile Phe 35	) Li ) T ) T  LECU: QUEN  Gly  Val 20  Asn	ENGT: YPE: OPOLO LE T CE D Trp 5 Val	H: OGY: YPE: YPE: ESCR Ser Pro	IPTI Glu Val Cys	ON: Ser	SEQ Leu Glu 25 Thr	amine line Prot ID N Lys 10 Thr	o ac ar ein O: Gly His	id 27: Ser Pro Arg	Glu Glu Cys 45	Leu 30	15 Thr Gly	Ser Cys
35	Asn 1 Arg Gln	(ii) (xi) Thr Pro Arg Asn 50	(A) (B) (D) MO: SEC Lys Ile Phe 35 Asp	) Li ) To ) To LECU: QUENC Gly Val 20 Asn	ENGT: YPE: OPOLO LE T CE D Trp 5 Val Pro	H: OGY: YPE: YPE: ESCR Ser Pro Pro	IPTIC Glu Val Cys Glu 55	ON: : Val Ser Val 40 Cys	SEQ Leu Glu 25 Thr	amineline. Prote ID No Lys 10 Thr Leu Pro	o accar ein O: Gly His	27: Ser Pro Arg	Glu Glu Cys 45 Glu	Leu 30 Gly	Thr Gly Asn	Ser Cys Val

5		(B) T	ENGTH: YPE: OPOLOGY:			amino aci o acid ar	ds	
	(ii)	MOLECUI	LE TYPE:		Prot	ein		
	(xi)	SEQUEN	CE DESCRI	PTION: S	SEQ ID N	0: 28:		
10	Ser Glu 1	Val Leu	Lys Gly 5	Ser Glu	Cys Lys 10	Pro Arg	Pro Ile	Val Val 15
15	Pro Val	Ser Glu 20	Thr His	Pro Glu	Leu Thr 25	Ser Gln	Arg Phe 30	Asn Pro
15	Pro Cys	Val Thr 35	Leu Met	Arg Cys 40	Gly Gly	Cys Cys	Asn Asp 45	Glu Ser
20	Leu Glu 50	Cys Val	Pro Thr	Glu Glu 55	Val Asn	Val Thr 60	Met Glu	Leu Leu
	Gly Ala 65	Ser Gly	Ser Gly 70	Ser Asn	Gly Met	Gln Arg 75	Leu Ser	Phe Val 80
25	Glu His	Lys Lys	Cys Asp 85	Cys Arg	Pro Arg 90	Phe Thr	Thr Thr	Pro Pro 95
2.0	Thr Thr	Thr Arg	Pro Pro	Arg Arg	Arg Arg 105			
30	(2) INF	ORMATION	FOR SEQ	ID NO:	29:			
	(i)	SEQUEN	CE CHARAG	CTERISTI	CS:			
35		(B) T	ENGTH: YPE: OPOLOGY:			amino ac: o acid ar	ids	
40	(ii)	MOLECU	LE TYPE:		Prot	ein		
40	(xi)	SEQUEN	CE DESCR	IPTION:	SEQ ID N	0: 29:		
45	Gly Ser 1		Lys Pro 5			Val Pro	Val Ser	Glu Thr 15
40	His Pro	Glu Leu 20	Thr Ser	Gln Arg	Phe Asr. 25	Pro Pro	Cys Val 30	Thr Leu
50	Met Arg	Cys Gly 35	Gly Cys	Cys Asn 40	Asp Glu	Ser Leu	Glu Cys 45	Val Pro
	Thr Glu 50	Glu Val	Asn Val	Thr Met 55	Glu Leu	Leu Gly 60	Ala Ser	Gly Ser
55	Gly Ser	Asn Gly	Met Gln	Arg Leu	Ser Pho	val Glu	His Lys	Lys Cys

	(2)	INFO	ORMA'I	CION	FOR	SEQ	ID N	10:	30:							
-		( <u>i</u> )	SEC	UENC	CE CI	IARAC	TERI	STIC	CS:							
5			(A) (B) (D)	TY	ENGTH PE: POLO					121 a amino linea	aci		lds			
10		(ii)	MOI	LECUI	LE TY	PE:			:	Prote	∍in					
		(xi)	SEÇ	QUENC	CE DE	ESCRI	PTIC	ON: S	SEQ	ID NO	): :	30:				
15	Asn 1	Asp	Ser	Pro	Pro 5	Ser	Thr	Asn	Asp	Trp 10	Met	Arg	Thr	Leu	Asp 15	Lys
	Ser	Gly	Cys	Lys 20	Pro	Arg	Asp	Thr	Val 25	Val	Tyr	Leu	Gly	Glu 30	Glu	Tyr
20	Pro	Glu	Ser 35	Thr	Asn	Leu	Gln	Tyr 40	Asn	Pro	Arg	Cys	Val 45	Thr	Val	Lys
25	Arg	Cys 50	Ser	Gly	Суѕ	Cys	Asn 55	Gly	Asp	Gly	Gln	Ile 60	Cys	Thr	Ala	Val
20	Glu 65	Thr	Arg	Asn	Thr	Thr 70	Val	Thr	Val	Ser	Val 75	Thr	Gly	Val	Ser	Ser 80
30	Ser	Ser	Gly	Thr	Asn 85	Ser	Gly	Val	Ser	Thr 90	Asn	Leu	Gln	Arg	11e 95	Ser
	Val	Thr	Glu	His 100	Thr	Lys	Cys	Asp	Cys 105	Ile	Gly	Arg	Thr	Thr 110	Thr	Thr
35	Pro	Thr	Thr 115	Thr	Arg	Glu	Pro	Arg 120	Arg							
	(2)	INF	ORMA!	rion	FOR	SEQ	ID 1	NO:	31:							
40		(i)	SEC	QUEN	CE CI	HARA(	CTER	ISTI	CS:							
			(A) (B) (D)	) T	ENGT: YPE: OPOL					116 amin amin line	o ac		ids			
45		(ii)	MO:	LECU:	LE T	YPE:				Prot	ein					
		(xi)	SE	QUEN	CE D	ESCR.	IPTI:	ON: :	SEQ	N CI	0:	31:				
50	Ser 1	Thr	Asn	Asp	Trp 5	Met	Arg	Thr	Leu	Asp 10	Lys	Ser	Gly	Cys	Lys 15	Pro
E.E	Arg	Asp	Thr	Val 20	Val	Tyr	Leu	Gly	Glu 25	Glu	Tyr	Pro	Glu	Ser 30	Thr	Asn
55	₫ eu	ci.		Acr	r · ·	. · ·	~·		- b		٠	*	*			• .

	65					70					75					80
r	Ser	Gly	Val	Ser	Thr 85	Asn	Leu	Gln	Arg	Ile 90	Ser	Val	Thr	Glu	His 95	Thr
5	Lys	Cys	Asp	Cys 100	Ile	Gly	Arg	Thr	Tnr 105	Thr	Thr	Pro	Thr	Thr 110	Thr	Arg
10	Glu	Pro	Arg 115	Arg												
	(2)	INFO	ORMA'	rion	FOR	SEQ	ID 1	10:	32:							
15		(i)	SE	QUENC	CE C	HARA(	CTER	ISTIC	cs:							
13			(A) (B) (D)	T	ENGTI PE: OPOLO					111 a amino linea	aci		ids			
20		(ii)	MOI	LECUI	LE T	YPE:				Prote	ein					
		(xi)	SE(	QUEN	CE DE	ESCRI	IPTIO	ON: S	SEQ	ID NO	): (	32:				
25	Met 1	Arg	Thr	Leu	Asp 5	Lys	Ser	Gly	Cys	Lys 10	Pro	Arg	Asp	Thr	Val 15	Val
	Tyr	Leu	Gly	Glu 20	Glu	Tyr	Pro	Glu	Ser 25	Thr	Asn	Leu	Gln	Tyr 30	Asn	Pro
30	Arg	Cys	Val 35	Thr	Val	Lys	Arg	Cys 40	Ser	Gly	Cys	Cys	Asn 45	Gly	Asp	Gly
35	Gln	Ile 50	Cys	Thr	Ala	Val	Glu 55	Thr	Arg	Asn	Thr	Thr 60	Val	Thr	Val	Ser
	Val 65	Thr	Gly	Val	Ser	Ser 70	Ser	Ser	Gly	Thr	Asn 75	Ser	Gly	Val	Ser	Thr 80
40	Asn	Leu	Gln	Arg	Ile 85	Ser	Val	Thr	Glu	His 90	Thr	Lys	Cys	Asp	Cys 95	Ile
	Gly	Arg	Thr	Thr 100	Thr	Thr	Pro	Thr	Thr 105	Thr	Arg	Glu	Pro	Arg 110	Arg	
45	(2)	INF	ORMA'	TION	FOR	SEQ	ID :	NO:	33:							
		( <u> </u>	SE:	QUEN	CE C	HARA(	CTER	ISTI	CS:							
50			•	) L: ) T:						106 . amin line	o ac		ids			
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55		(xi)	SE	QUEN	CE D	ESCR	IFTI	ON:	SEQ	n cı	0:	33:				

	Lys	Arg	Cys 35	Ser	Gly	Cys	Cys	Asn 40	Gly	Asp	Gly	Gln	Ile 45	Суѕ	Thr	Ala
5	Val	Glu 50	Thr	Arg	Asn	Thr	Thr 55	Val	Thr	Val	Ser	Val 60	Thr	Gly	Val	Ser
	Ser 65	Ser	Ser	Gly	Thr	Asn 70	Ser	Gly	Val	Ser	Thr 75	Asn	Leu	Gln	Arg	Ile 80
10	Ser	Val	Thr	Glu	His 85	Thr	Lys	Cys	Asp	Cys 90	Ile	Gly	Arg	Thr	Thr 95	Thr
15	Thr	Pro	Thr	Thr 100	Thr	Arg	Glu	Pro	Arg 105	Arg						
10																
	(2)	INFO	ORMAT	NOI	FOR	SEQ	ID 1	10:	34:							
20		(i)	SEÇ	QUENC	CE C	IARA(	CTER	ISTIC	CS:							
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25		(ii)	MOI	LECUI	LE T	YPE:			I	Prot€	ein					
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30	Pro 1	Val	Ser	Gln	Phe 5	Asp	Gly	Pro	Ser	His 10	Gln	Lys	Lys	Val	Val 15	Pro
	Trp	Ile	Asp	Val 20	Tyr	Thr	Arg	Ala	Thr 25	Cys	Gln	Pro	Arg	Glu 30	Val	Val
<b>3</b> 5	Val	Pro	Leu 35	Ser	Met	Glu	Leu	Met 40	Gly	Asn	Val	Val	Lys 45	Gln	Leu	Val
40	Pro	Ser 50	Cys	Val	Thr	Val	Gln 55	Arg	Cys	Gly	Gly	Cys 60	Cys	Pro	Asp	Asp
40	Gly 65	Leu	Glu	Cys	Val	Pro 70	Thr	Gly	Gln	His	Gln 75	Val	Arg	Met	Gln	Ile 80
45	Leu	Met	Ile	Gln	Tyr 85	Pro	Ser	Ser	Gln	Leu 90	Gly	Glu	Met	Ser	Leu 95	Glu
	Glu	His	Ser	31n 100	Cys	Glu	Cys	Arg	Pro 105	Lys	Lys	Lys	Glu	Ser 110	Ala	Val
50	Lys	Pro	Asp 115	Ser	Pro	Arg	Ile	Leu 120	Cys	Pro	Pro	Cys	Thr 125	Gln	Arg	Arg
55	Gln	Arg 130	Pro	Asp	Pro	Arg	Thr 135	Cys	Arg	Cys	Arg	Cys 140	Arg	Λrg	Arg	Arg
JJ	Phe	I.evi	Hic	عبب	dir.	?·	Ž mir	a ·	• .	<i>~</i> :	• 0	•		•		

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10		(xi)	SE	QUEN	CE DE	ESCR	IPTIC	: NC	SEQ	ID NO	): :	35:				
	Pro 1	Val	Ser	Gln	Pro 5	Asp	Ala	Pro	Gly	His 10	Gln	Arg	Lys	Val	Val 15	Ser
15	Trp	Ile	Asp	Val 20	Tyr	Thr	Arg	Ala	Thr 25	Cys	Gln	Pro	Arg	Glu 30	Val	Val
20	Val	Pro	Leu 35	Thr	Val	Glu	Leu	Met 40	Gly	Thr	Val	Ala	Lys 45	Gln	Leu	Val
_ `	Pro	Ser 50	Cys	Val	Thr	Val	Gl դ 55	Arg	Суѕ	Gly	Gly	Суз 60	Суз	Pro	Asp	Asp
25	Gly 65	Leu	Glu	Cys	Val	Pro 70	Thr	Gly	Gln	His	Gln 75	Val	Arg	Met	Gln	Ile 80
	Leu	Met	Ile	Arg	Tyr 85	Pro	Ser	Ser	Gln	Leu 90	Gly	Glu	Met	Ser	Leu 95	Glu
30	Glu	His	Ser	Gln 100	Cys	Glu	Cys	Arg	Pro 105	Lys	Lys	Asp	Ser	Ala 110	Val	Lys
35	Pro	Asp	Arg 115	Ala	Ala	Thr	Pro	His 120	His	Arg	Pro	Gln	Pro 125	Arg	Ser	Val
	Pro	Gly 130	Trp	Asp	Ser	Ala	Pro 135	Gly	Ala	Pro	Ser	Pro 140	Ala	Asp	Ile	Thr
40	His 145	Pro	Thr	Pro	Ala	Pro 150	Gly	Pro	Ser	Ala	His 155	Ala	Ala	Pro	Ser	Thr 160
	Thr	Ser	Ala	Leu	Thr 165	Pro	Gly	Pro	Ala	Ala 170	Ala	Ala	Ala	Asp	Ala 175	Ala
45	Ala	Ser	Ser	Val 180	Ala	Lys	Gly	Gly	Ala 185							
50	(2)	INFO	ORMAT	LION	FOR	SEQ	ID 1	10:	36:							
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	Trp	Ile	Asp	Val 20	Tyr	Thr	Arg	Ala	Thr 25	Cys	Gln	Pro	Arg	Glu 30	Val	Val
5	Val	Pro	Leu 35	Thr	Val	Glu	Leu	Met 40	Gly	Thr	Val	Ala	Lys 45	Gln	Leu	Val
1.0	Pro	Ser 50	Cys	Val	Thr	Val	Gln 55	Arg	Суз	Gly	Gly	Cys 60	Суѕ	Pro	Asp	Asp
10	Gly 65	Leu	Glu	Cys	Val	Pro 70	Thr	Gly	Gln	His	Gln 75	Val	Arg	Met	Gln	Ile 80
15	Leu	Met	Ile	Arg	Tyr 85	Pro	Ser	Ser	Gln	Leu 90	Gly	Glu	Met	Ser	Leu 95	Glu
	Glu	His	Ser	Gln 100	Cys	Glu	Cys	Arg	Pro 105	Lys	Lys	Lys	Asp	Ser 110	Ala	Val
20	Lys	Gln	Asp 115	Arg	Ala	Ala	Thr	Pro 120	His	His	Arg	Pro	Gln 125	Pro	Arg	Ser
25	Val	Pro 130	Gly	Trp	Asp	Ser	Ala 135	Pro	Gly	Ala	Pro	Ser 1 <b>4</b> 0	Pro	Ala	Asp	Ile
23	Thr 145	Gln	Ser	His	Ser	Ser 150	Pro	Arg	Pro	Leu	Cys 155	Pro	Arg	Cys	Thr	Gln 160
30	His	His	Gln	Cys	Pro 165	Asp	Pro	Arg	Thr	Cys 170	Arg	Cys	Arg	Cys	Arg 175	Arg
	Arg	Ser	Phe	Leu 180	Arg	Cys	Gln	Gly	Arg 185	Gly	Leu	Glu	Leu	Asn 190	Pro	Asp
35	Thr	Cys	Arg 195	Cys	Arg	Lys	Leu	Arg 200	Arg							
	(2)	INF	ORMA:	rion	FOR	SEQ	ID	:01/	37:							
40		(i)	SEÇ	QUEN	CE CI	HARA	CTER:	ISTI	CS:							
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30	1	PIO	Arg	GIU	A1a 5	Pro	AIG	Ald	Ата	10	Ala	Phe	Glu	Ser	15	Leu
55	Asp	Leu	Ser	Asp 20	Ala	Glu	Pro	Asp	Ala 25	Gly	Glu	Ala	Thr	Ala 30	Tyr	Ala
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	Leu 65	Arg	Lys	Gly	Gly	Trp 70	Gln	His	Asn	Arg	Glu 75	Gln	Ala	Asn	Leu	Asr 80
5	Ser	Arg	Thr	Glu	Glu 85	Thr	Ile	Lys	Phe	Ala 90	Ala	Ala	His	Tyr	Asn 95	Thr
	Glu	Ile	Leu	Lys 100	Ser	Ile	Asp	Asn	Glu 105	Trp	Arg	Lys	Thr	Gln 110	Cys	Met
10	Pro	Arg	Glu 115	Val	Cys	Ile	Asp	Val 120	Gly	Lys	Glu	Phe	Gly 125	Val	Ala	Thr
15	Asn	Thr 130	Phe	Phe	Lys	Pro	Pro 135	Суз	Val	Ser	Val	Tyr 140	Arg	Cys	Gly	Gly
13	Cys 145	Cys	Asn	Ser	Glu	Gly 150	Leu	Gln	Cys	Met	Asn 155	Thr	Ser	Thr	Ser	Tyr 160
20	Leu	Ser	Lys	Thr	Leu 165	Phe	Glu	Ile	Thr	Val 170	Pro	Leu	Ser	Gln	Gly 175	Pro
	Lys	Pro	Val	Thr 180	Ile	Ser	Phe	Ala	Asn 185	His	Thr	Ser	Cys	Arg 190	Cys	Met
25	Ser	Lys	Leu 195	Asp	Val	Tyr	Arg	Gln 200	Val	His	Ser	Ile	Ile 205	Arg	Arg	Ser
30	Leu	Pro 210	Ala	Thr	Leu	Pro	Gln 215	Cys	Gln	Ala	Ala	Asn 220	Lys	Thr	Cys	Pro
	Thr 225	Asn	Tyr	Met	Trp	Asn 230	Asn	His	Ile	Суѕ	Arg 235	Суѕ	Leu	Ala	Gln	Glu 240
35	Asp	Phe	Met	Phe	Ser 245	Ser	Asp	Ala	Gly	Asp 250	Asp	Ser	Thr	Asp	Gly 255	Ph∈
	His	Asp	Ile	Cys 260	Gly	Pro	Asn	Lys	Glu 265	Leu	Asp	Glu	Glu	Thr 270	Cys	Glr
40	Cys	Val	Cys 275	Arg	Ala	Gly	Leu	Arg 280	Pro	Ala	Ser	Суз	Gly 285	Pro	His	Lys
45	Glu	Leu 290	Asp	Arg	Asn		Cys 295		Cys	Val	_	Lys 300		Lys	Leu	Ph∈
	Pro 305	Ser	Gln	Cys	Gly	Ala 310	Asn	Arg	Glu	Phe	Asp 315	Glu	Asn	Thr	Cys	Glr 320
50	Cys	Val	Cys	Lys	Arg 325	Thr	Cys	Pro	Arg	Asn 330	Gln	Pro	Leu	Āsn	Pro 335	Gly
	Lys	Cys	Ala	Cys 340	Glu	Cys	Thr	Glu	Ser 345	Pro	Gln	Lys	Cys	Leu 350	Leu	Lys
55	Gly	Lys	Lys	Phe	His	His	Gln	Thr	Cys	Ser	Суз	Tyr	Arg	Arg	Pro	Суз

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5			(A) (B) (D)	TY	ENGTI (PE: OPOLO					133 a amino linea	ac		ids			
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		(xi)	SEÇ	QUENC	CE DE	ESCRI	PTIC	ON: S	SEQ	ID N	): :	38:				
15	Met 1	Lys	Leu	Leu	Val 5	Gly	Ile	Leu	Val	Ala 10	Val	Cys	Leu	His	Gln 15	Tyr
	Leu	Leu	Asn	Ala 20	Asp	Ser	Asn	Thr	Lys 25	Gly	Trp	Ser	Glu	Val 30	Leu	Lys
20	Gly	Ser	Glu 35	Суѕ	Lys	Pro	Arg	Pro 40	Ile	Val	Val	Pro	Val 45	Ser	Glu	Thr
25	His	Pro 50	Glu	Leu	Thr	Ser	Gln 55	Arg	Phe	Asn	Pro	Pro 60	Cys	Val	Thr	Leu
	Met 65	Arg	Cys	Gly	Gly	Cys 70	Суѕ	Asn	Asp	Glu	Ser 75	Leu	Glu	Суз	Val	Pro 80
30	Thr	Glu	Glu	Val	Asn 85	Val	Thr	Met	Glu	Leu 90	Leu	Gly	Ala	Ser	Gly 95	Ser
	Gly	Ser	Asn	Gly 100	Met	Gln	Arg	Leu	Ser 105	Phe	Val	Glu	His	Lys 110	Lys	Cys
35	Asp	Cys	Arg 115	Pro	Arg	Phe	Thr	Thr 120	Thr	Pro	Pro	Thr	Thr 125	Thr	Arg	Pro
40	Pro	Arg 130	Arg	Arg	Arg											
	(2)	INF	ORMAT	пои	FOR	SEQ	ID 1	NO:	39:							
45		(i)	SEÇ	QUENC	CE CI	HARA	CTER	ISTIC	CS:							
43			(A) (B) (D)	T	ENGTI YPE: DPOLO					148 amino amino line	o ac:		ids			
50		(ii)	MOI	LECUI	LE T	YPE:				Prot	ein					
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55	Met 1	Lys	Leu	Thr	Ala 5	Thr	Leu	Gln	Val	Val 10	Val	Ala	Leu	Leu	Ile 15	Cys

Arg Asp Thr Val Val Tyr Leu Gly Glu Glu Tyr Pro Glu Ser Thr Asn 50 55 Leu Gln Tyr Asn Pro Arg Cys Val Thr Val Lys Arg Cys Ser Gly Cys 5 Cys Asn Gly Asp Gly Gln Ile Cys Thr Ala Val Glu Thr Arg Asn Thr 10 Thr Val Thr Val Ser Val Thr Gly Val Ser Ser Ser Ser Gly Thr Asn Ser Gly Val Ser Thr Asn Leu Gln Arg Ile Ser Val Thr Glu His Thr 15 Lys Cys Asp Cys Ile Gly Arg Thr Thr Thr Thr Pro Thr Thr Thr Arg 135 Glu Pro Arg Arg 20 145 (2) INFORMATION FOR SEQ ID NO: 40: (i) SEQUENCE CHARACTERISTICS: 25 (A) LENGTH: 26 amino acids amino acid (B) TYPE: (D) TOPOLOGY: linear 30 (ii) MOLECULE TYPE: Protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40: Met Asn Phe Leu Leu Ser Trp Val His Trp Ser Leu Ala Leu Leu Leu 35 10 Tyr Leu His His Ala Lys Trp Ser Gln Ala 40 (2) INFORMATION FOR SEQ ID NO: 41: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: (B) TYPE: 20 base pairs ' 45 nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41: 50

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GCAGAGCTCG TTTAGTGAAC

## Claims

- A truncated VRP subunit having a deletion of at least one of the amino acid residues N-terminal to the first cysteine
   of the core sequence of said subunit.
  - 2. The truncated VRP subunit of claim 1 wherein the VRP is a human VRP.
- 3. The truncated VRP subunit of claim 1 wherein said VRP is selected from the group consisting of VEGF-B, VRF-2, VEGF-C, PlGF, VEGF-3, poxvirus ORF-1, and poxvirus ORF-2.
- 4. The truncated VRP subunit of claim 1 wherein said VRP 15 is VEGF-B.
  - 5. The truncated VRP subunit of claim 1 wherein said VRP subunit comprises an amino acid sequence of Figure 2.
- 20 6. The truncated VRP subunit of claim 1 wherein the amino acid residues N-terminal to the first cysteine of the core sequence of said subunit are deleted.
- 7. The truncated VRP subunit of claim 1 wherein the 25 amino acid sequence N-terminal to said core sequence comprises 2 to 5 amino acid residues.
- 8. The truncated VRP subunit of claim 7 wherein said 2 to 5 amino acid residues comprise 2 to 5 of the consecutive 30 amino acid residues immediately N-terminal to the first cysteine of the core sequence of said VRP subunit.

35 6 to 13 amino acid residues.

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- 10. The truncated VRP subunit of claim 1 wherein said 6 to 10 amino acid residues comprise 6 to 10 of the consecutive amino acid residues immediately N-terminal to the first cysteine of the core sequence of said VRP subunit.
  - 11. The truncated VRP subunit of claim 1 wherein the amino acid sequence N-terminal to said core sequence comprises 11 to 20 amino acid residues.

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12. The truncated VRP subunit of claim 1 wherein said 11 to 20 amino acid residues comprise 11 to 20 of the consecutive amino acid residues immediately N-terminal to the first cysteine of the core sequence of said VRP subunit.

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13. The truncated VRP subunit according to claim 1, further comprising at the N-terminus of said truncated VRP subunit, the first one or two amino acid residues of the mature non-truncated VRP subunit.

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- 14. A truncated VRP comprising two VRP subunits of claim 13.
- 15. A truncated VRP comprising two VRP subunits of claim
  25 1, wherein said two VRP subunits have the same amino acid
  sequence.
  - 16. A truncated VRP heterodimer comprising

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- a first subunit comprising a truncated VRP subunit of 30 claim 1; and
  - a second subunit comprising a subunit selected from the

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- 17. A nucleic acid molecule coding for a truncated VRP subunit of claim 1.
- 5 18. The nucleic acid molecule of claim 17 wherein the nucleic acid molecule is a DNA molecule.
  - 19. The nucleic acid molecule of claim 17 wherein the nucleic acid molecule is an RNA molecule.

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- 20. A recombinant DNA vector comprising the nucleic acid molecule of claim 17.
- 21. A recombinant DNA expression vector comprising a 15 nucleic acid molecule of claim 17.
- 22. The recombinant DNA expression vector of claim 21 wherein said nucleic acid molecule is operably linked at the 5' end of said nucleic acid molecule to a DNA sequence that codes 20 for a signal peptide.
  - 23. The recombinant DNA expression vector of claim 22 wherein said signal peptide is selected from the group consisting of VEGF signal peptide, VEGF-B signal peptide, VRF-2 signal peptide, VEGF-C signal peptide, VEGF-3 signal peptide, and PlGF signal peptide.
- 24. The recombinant DNA expression vector of claim 22 wherein said signal peptide is selected from the group 30 consisting of poxvirus ORF-1 signal peptide, and poxvirus ORF-2 signal peptide.

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26. The recombinant DNA expression vector of claim 22 wherein said DNA sequence coding for said signal peptide is operably linked at the 3' end of said DNA sequence to DNA coding for the first amino acid residue of the mature non-truncated VRP subunit and wherein the 3' end of said DNA coding for said residue is operably linked to said nucleic acid molecule coding for said truncated VRP subunit.

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- 27. The recombinant DNA expression vector of claim 22 wherein said DNA sequence coding for said signal peptide is operably linked at the 3' end of said DNA sequence to DNA coding for the first two amino acid residues of the mature non-truncated VRP subunits and wherein the 3' end of said DNA coding for said two residues is operably linked to said nucleic acid molecule coding for said truncated VRP subunit.
  - 28. The recombinant DNA expression vector of claim 22 wherein said nucleic acid molecule is operably linked to control sequences operable in a host cell transformed with said vector.
    - 29. A transformed or transfected host cell comprising the recombinant DNA expression vector of claim 21.
- 25 30. A transformed or transfected host cell comprising the recombinant DNA expression vector of claim 22.
  - 31. A transformed or transfected host cell comprising the recombinant DNA expression vector of claim 26.
  - 32. A delivery vector comprising a nucleic acid molecule

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- 33. A delivery vector of claim 32, wherein said delivery vector is a viral delivery vector.
- 34. An adenovirus vector comprising the nucleic acid molecule of claim 17.
- 35. The adenovirus vector of claim 34 wherein said nucleic acid molecule is operably linked at the 5' end of said nucleic acid molecule to a DNA sequence that codes for a signal 10 peptide.
  - 36. The adenovirus vector of claim 35 wherein said signal peptide is selected from the group consisting of VEGF signal peptide, VEGF-B signal peptide, VRF-2 signal peptide, VEGF-C signal peptide, and PIGF signal peptide.
    - 37. The adenovirus vector of claim 35 wherein said signal peptide is selected from the group consisting of poxvirus ORF-1 signal peptide, and poxvirus ORF-2 signal peptide.

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- 38. The adenovirus vector of claim 35 wherein said signal peptide is VEGF-B signal peptide.
- 39. The adenovirus vector of claim 35 wherein said DNA sequence coding for said signal peptide is operably linked at the 3' end of said DNA sequence to DNA coding for the first amino acid residue of the mature non-truncated VRP subunit, and wherein the 3' end of said DNA coding for said residue is operably linked to said nucleic acid molecule coding for said truncated VRP subunit.

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- 40. A filtered injectable adenovirus vector preparation, comprising: a recombinant adenoviral vector, said vector containing no wild-type virus and comprising:
- a partial adenoviral sequence from which the  ${\tt ElA/ElB}$  genes base been deleted, and
  - a transgene coding for a truncated VRP subunit of claim 1, driven by a promoter flanked by the partial adenoviral sequence; and
    - a pharmaceutically acceptable carrier.

- 41. The preparation of claim 40 wherein said adenovirus vector has been filtered through a 30 micron filter.
- 42. The injectable adenoviral vector preparation according to claim 40 wherein said promoter is selected from the group consisting of a CMV promoter, a ventricular myocytespecific promoter, and a myosin heavy chain promoter.
- 43. A method of producing a truncated VRP polypeptide comprising growing, under suitable conditions, a host cell transformed or transfected with the recombinant DNA expression vector of claim 21 in a manner allowing expression of said polypeptide, and isolating said polypeptide from the host cell.
- 25 44. A pharmaceutical composition comprising a VRP comprising at least one truncated VRP subunit of claim 1, in a suitable carrier.
- 45. A method of stimulating blood vessel formation 30 comprising administering to a patient a pharmaceutical composition comprising a truncated VRP comprising at least one

46. A method of stimulating endothelial cell growth or cell migration in vitro comprising treating said endothelial cells with a truncated VRP comprising at least one truncated VRP subunit of claim 1, in a suitable carrier.

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- 47. A method of treating a patient suffering from a heart disease comprising administering to said patient a nucleic acid molecule coding for at least one truncated VRP subunit of claim 1, said nucleic acid molecule capable of expressing the truncated VRP subunit in said patient.
- 48. A method of stimulating angiogenesis in a patient comprising administering a therapeutically effective amount of a pharmaceutical composition comprising a truncated VRP comprising at least one truncated VRP subunit of claim 1, in a suitable carrier.
- 49. The method of claim 48 further comprising a therapeutically suitable delivery system for said pharmaceutical composition.
  - 50. The method of claim 48 further comprising administering a potentiating agent that potentiates the angiogenic effect of said truncated VRP.

- 51. The method of claim 50, wherein said potentiating agent is an angiogenic FGF.
- 52. The method of claim 51, wherein said potentiating 30 agent is selected from the group consisting of FGF-1, FGF-2, FGF-4, FGF-5, and FGF-6.

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and one or more potentiating agents in a pharmaceutically acceptable carrier.

- 54. The pharmaceutical composition of claim 53 wherein said potentiating agent is an angiogenic FGF.
- 55. The pharmaceutical composition of claim 54, wherein said potentiating agent is selected from the group consisting of FGF-1, FGF-2, FGF-4, FGF-5, and FGF-6, in a pharmaceutically acceptable carrier.
- 56. A method of treating a patient suffering from an ischemic condition comprising administering a therapeutic amount of a pharmaceutical composition comprising a truncated VRP comprising at least one truncated VRP subunit of claim 1, in a suitable carrier.
- 57. The method of claim 56 further comprising administering an agent that potentiates the therapeutic effect 20 of said truncated VRP subunit.
  - 58. The method of claim 57 wherein said potentiating agent is selected from the group consisting of FGF-1, FGF-2, FGF-4, FGF-5, and FGF-6.

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59. The method of claim 56 wherein said ischemic condition is selected from the group consisting of: cardiac infarction, chronic coronary ischemia, chronic lower limb ischemia, stroke, and peripheral vascular disease.

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60. A method for treating a patient suffering from a

comprising at least one truncated VRP subunit according to claim 1, in a suitable carrier.

- 61. A method of increasing vascular permeability comprising administering a therapeutic amount of a pharmaceutical composition comprising a truncated VRP comprising at least one truncated VRP subunit according to claim 1, in a suitable carrier.
- 10 62. A method of stimulating angiogenesis in a patient comprising delivering a delivery vector to the myocardium of the patient by intracoronary injection directly into one or both coronary arteries, said vector comprising a nucleic acid molecule coding for at least one truncated VRP subunit according to claim 1, wherein said vector is capable of expressing the truncated VRP subunit in the myocardium.
  - 63. The method of claim 62, wherein said delivery vector is a replication-deficient adenovirus vector.

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- 64. A method for stimulating coronary collateral vessel development in a patient having myocardial ischemia, comprising delivering a delivery vector to the myocardium of the patient by intracoronary injection directly into one or both coronary arteries, said vector comprising a nucleic acid molecule coding for a truncated VRP subunit and capable of expressing the truncated VRP subunit in the myocardium, thereby promoting coronary collateral vessel development.
- 30 65. The method of claim 64, wherein said delivery vector is a replication-deficient adenovirus vector.

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delivering a delivery vector to the peripheral vascular system of the patient by intra-femoral artery injection directly into one or both femoral arteries, said vector comprising a transgene coding for a truncated VRP subunit, and capable of expressing the truncated VRP subunit in the peripheral vascular system, thereby promoting peripheral vascular development.

67. The method of claim 66, wherein said delivery vector is a replication-deficient adenovirus vector.

## Figure 1

hvegFC	mh.lgffsvacsllaaallpGPREAPAAAAFESGLDLSDAEP
hVEGFB hVRF2 hVFGFC	msplirtiivaliqiartqaPVSQFDGPSHQKKVVPWIDVYTHAT msplirtillaaliqlapaqaPVSQPDAPGHQRKVVSWIDVYTRAT naqealayaSKDLEEGLESVSSVDELKTVLYPEYWHWYRQDLERGSWQHNREQAXLNSPTEETIKFAAAHTHTEILFSTTWENTQ
hPlGF	mpvmrlfpcflqllaglalpAVPPQQWALSAGNGSSEVEVVPFQEVKGRSY
hVEGF3 DVORF1	mrrcrisgrppappgvpaqaPVSQPDAPGHQRKVVSWIDVITRAI MKLLVGILVAVCLHQYLINADSNTKGWSEVIKGSE
pvorf2	MKLTRITEQVVVALLICHYNLPECVSQSNDSPPSTNDWMRTLDRSG
hvegfb	SSEEVVVPLSMELMGRVVFQLVPSGVTVQPQSGCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQ
hVRF2	COPREVVVPLTVELMGTVAKQLVPSCVTVQRCGGCCPDDGLECVPTGGHQVRMQILMIRYPSSQLGEMSLEEHSQCEC CMDDEGGTTRGGEGGGAATMTEFKDDGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
nvege hplan	CRETEVILD VOLDE OVALINITE IN TENTO COLOR CONDIGUES CONTRACTOR DE LA CONTRACTOR DE LA CONTRACTOR DE C
hVEGF3	COPREVIVELIMETVARQLVPSCVTVQPCGCCCPDDGLECVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEC
pv0RF1	CKPRPIVVPVSETHPELTSQRFNPPCVTLMRCGGCCNDESLECVPTERVNVTMELLGASGSGSNGMQRLSFVEHKKCDC
pv0RF2	<u>CKPPDTVVYLGEEYPESTHLQYHPRCVTVKPGSGCCHGDGQICTAVETRHTTVTVSVTGVSSSSGTNSGVSTHLQRISVTEHTKCDC</u>
hvegfb	R PYYKESAVK PDSPP II. CPPCTQRPQP PDPRICPCRCRRRFIHCQGFGLELNPDICRCRKPRK
hVRF2	RPKKDSAVKPDRAATPHHRPQPRSVPGWDSAPGAPSPADITHPTPAPGPSAHAAPSTTSALTPGPAAAAADAASSVAKGGA
hvegfc	MSYLDVYRQVHSIIRRSLPATI,PQCQAANKTCPTNYMWNNHICRCLAÇEDFMFSSDAGDDSTDGFHDICGPNKELDEETCQCVCRAG
hPlGF	R PLI REKYNKPERRRPKGRGKRRREKQRPTDCHLCGDAVPRR
hVEGF3	RPEKKDSAVKODPAATPHHRPOPPSVPGWDSAPGAPSPADITOSHSSPRPLCPRCTOHHOCPDPRTCRCRCRRRSFIRCOGRGLELN
pv0RF1	RPRETITIPPITIRPPRRER
PVORF2	IGRITITITITITEPRR
hvegfc	LRPASCGPHKELDPINSCQCVCYNKLFPSQCGANREFDENTCQCVCYPTJPRNQPLNPGYCACECTESPQKCLLYGYRFHHQTGSYR
hvegf3	PUTCRCRKLRR
hvegyc	E PCTNRQKACEPGFSYSEEVCRCVPSYWKRPPQMS

IDVYTRAT YTRAT EAT

PVSQFDGPSHQKKVVPWIDVYTRAT PSHQKKVVPWIDVYTRAT

KVVPWIDVYIRAT FWIDVYTRAT

Figure 2a VEGF-B CQPREVVPLSMELHGHYVK\_LUV FSCYTVGRGGGGGPOT POUGLEGVFT AQHQVRMQILMIQYFGGGLGFMGT FTH NGGFO OQPREVVPLSMELMGNVVKQLV PSGVTVQRGGGGGPOUGLEGVPT GQHQVRMQILMIQYPS SQLGEMSLEBHSQGBG GQPREVVPLSMELMGNVVKQLVPSGVTVQRGGGGPOUGLEGVPT GQHQVRMQILMIQYPS SQLGEMSLEBHSQGBG CQPREVVPLSMELMGNVVKQLVPSGVTVQRGGGGPOUGLEGVPT GQHQVRMQILMIQYPS SQLGEMS LEHSQGBG CQPREVVPLSMELMGNVVKQLVPSGVTVQRGGGGCPOUGLEGVPT GQHQVRMQILMIQYPS SQLGEMS LEHSQGBG GQPREVVPLSMELMGNVVKQLVPSGVTVQRGGGGCPOUGLEGVPT GQHQVRMQILMIQYPS SQLGEMS LEHSQGBG GQPREVVPLSMELMGNVVKQLVPSGVTVQRGGGGCPDDGLEGVPT GQHQVRMQILMIQYPS SQLGEMS LEHSQGBG ELPEKFESAVFDSPPTLOFFCTTPPGRFDPRTCPOFOFRFFTHOGGROLELNPDTCROPKFPK FLYKFESAVFDSPPTLOFFCT, PPLPFDPPTGROFOFPFFFUHOGSROLELNFDTCROPFFFK

500000

F/1 (1) (2) (3) (4) (5) (6)

ELEKKESAJVRPOSPRILCPPCTOPPOPRDPRICECROPPFFLHOGGRGLELNPDTCRORKPPY RPKKKESAVKPOSPRILCPPCTOPPOPPDPPTCRORCFRPPFLHOOGGPGLELNPDTCRORKPPK RPKKESAVKPOSPPILOPPCTORPOPPDPTCFCROFRPFIHOOGRGLELNPDTCRORKPPK

RPEKKESAVKPDSPFILCPPCTQPRQRPDPRTCPCPCPRBRPFLHCQGKGLELNPDTCRCRKPPK KLKEKESAVKPDSPFILCPPCTQPPQPPDPFTCPCPCFFRFFLHCQGRGLELNPDTCRCPKPPK

E7.1 (2) (3) (4) (6) (6) (6)

PVSQPDAPGHQRKVVSWIDVYTRAT
PGHQRKVVSWIDVYTRAT
KVVSWIDVYTRAT
IDVYTRAT

(£) (£) (£) (£)

Figure 2b VRF-2

COPPEVVVPLTVELMGTVAROLVPSGVTVPPGGGCPPDAGLGGVPTGGHQVPMQILMIRYPSSQFGEMSLEEHSGGLG	PPKKDSAVKPDRAATPHHPPPPRSVPGWDSAPGAESEADIIHPIPAPGPSAHAAPSITSALTPGFAAAADUALASSVARCCL
COPREVVPLTVELMGTVAROLVPSGVTVQRGGGGPDDGGLGGVPTGGHQVRMQILMIRYPSSQLGEMSLEEHSGGEG	RPKKDGAVFPDRAATPHHFPPPRSVPGWDSAPGAPSFRJITHITPAFGFSAHAAPSITSALTPGFAAAAAAAASSVARCGA
COPREVVVPLTVELMGTVARQLVPSGVTVQRGGGCPPDGSLEGVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSGGEG	RPKKDSAVKPDRAATPHHRPPPRSVPGWDSAPGAPSPADIIHPITPAFGPSAHAAPSITSALTPGFAAAAAAASSVAKGGA
COPREVVVPLTVELMGTVARQLVPSGVTVQRGGGGGPPDGSLEGVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSGGEG	RPKKDSAVKPDRAATPHHRPPPRSVPGWDSAPGAPSFADIIHPITPAFGPSAHAAPSITSALTPGFAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
F/L (1) (2) (3)	F/I (1) (2) (3) (4)

PGHQRKVVSWIEVYTRAT

KVVSWIDVYTRAT

(2) (2) (3) (4)

PVSQPDAPGHQRKVVSWICVYTRAT

IDVYTRAT

Figure 2c VEGF-3

REKKKUSAVKOOPRAJTPHIIPPOPRSVEGWOSAPGARSPADI TOSKSSPRDLOPPOTQHHQOPDPRTORORFESET ROGERGI BLIN REKKKUSAVKOORAAT PHIIRAQERSVPGWOSARGARSPADI TOSKSSPRPLOPPOTQHHQOPDPRTORORPHSKLLE OOGBGLEIN REKKKUSAVKQORAAT PHHREQPRSVPGWDSAPGAPSPADI TOSKSSPRPLOPPOTQHHQOPDPRTORORPPSFLP OOGPGLEIN REKKKUSAVKQORAAT PHHREQPRSVPGWUSAPGAPSPADI TOSKSSPRPLOPPOTQHHQOPDPRTORORPPSFLF OOGPGLEIN REKKKUSAVKQORAAT PHHREQPRSVPGWUGAPGAPSTADI TOOHO TERELOFROTQHHQOPDPRTORORPPRSFLF OOFF FOLEIN C\_PREVVPLTVELMSTVAKQLVESGVTVQRGGGCPDDGLEGVETGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEG GCPREVVPLTVELMSTVAKQLVPSGVTVQRGGGCPDDGLEGVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEG GQPREVVPLTVELMSTVAKQLVPSGVTVQRGGGCPDDGLEGVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEG GQPREVVVPLTVELMSTVAKQLVPSGVTVQRGGGCCPDDGLEGVPTGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEG GQPREVVVPLTVELMSTVAKQLVFSGVTVQRGGGCCPDDGLEGVFTGQHQVRMQILMIRYPSSQLGEMSLEEHSQCEG PDTCRCRKLRR PDTCRCRKLRR PUTCRCRKIRR PUTCRCRKLPR PUTCRCRKLRR EZ (2) (2) (3) (3) (3) E/L (2) (2) (3) (4) (4) (1) (2) (3) (4) Figure 2d VEGF-C

GPREAPAAAFESGLDLSDAEF	omsertryrsytetikstikstoksytruestyksyksionenytykysyksyksysykytresynyskteetikeralestyksionenkki hynteilksionenkkio ilksionenkky onenkkys	CHPREVCIDVGKERGVATNIFEKPROGOGOGNSEGLOGMNISTSYLSKTLFEITVPLSQGPKPVIISFANHTSGRO CMPREVTIDVGKERGVATNIFEKPPÖVSYYPÖGGOGNSEGLOGMNISTSYLSKTLFEITVPLSQGPKPVIISFANHTSÖRÖ CHFREVCIDVGKERGVATNIFEKFPÖVSYYPÖGGOGNSEGLOGMNISTSYLSKTLFEITVPLSQGPKRVIISFANHTSÖRÖ CMFREVCIDVGKERGVATNIFEKPPÖVSYYPÖGGOGNSEGLOGMNISTSYLSKTLFEITVPLSQGPKRVIISFANHTSÖRÖ CMFREVCIDVAKERGVATNIFEKPPÖVSYYPÖGGCONSEGLOGMNISTSYLSKTLFEITVPLSQGPKRVIISFANHTSÖRÖ	MSKLLVVRÐVHSTIRRSIRATLEGGGAANFTOFTNYMWNRIGFOFTINGEDEMESSBASDOSTIGEHDIGGENKELLEETGGVORAS MSFLDVYRGVHSIIRRSIRATLEGGAANFTOFTNYMWNRIOFOLAGEDFMESSBASDOSTDGFHDIGGENKEL DETGLVYGRAS MSFLDVYRGVHSIIRRSIRATLEGGAANFTOFTNYMWNRIOFOLAGEDFMESSBASDOSTDGFHDIGGENKEL FETGLVYGRAS MSKLDVYRGVHSIIRRSLPATLEGGAANFTOFTNYMWNRIOFOLAGEDFWESSBASDDSTDGFHDIGGENKELDE ETGLVYGKAS	LAPASOGEHKELDPRISOLTVORMELE PSQUGANBEFDENTOLGOVOKETOPENDELNPGKOAGECTES PQROLLKGRREHH, POCOTR LAFAS GEFHKELDPRISOLTVORNET EPSQUGANBEFDENTOLGOVORET TEPRNDELNFSKOAGECTES PQROLLK HEFFRELT OVR LEFASOGEHKELDPRISOLVORNELFPSQUGANBEFDENTOLGOVORETOPRNDELNFGKOAGECTES PQROLLK SPETRH JT COOTR FPASOGEHKELDPRISOLOVORNELFPSQUGANPEFDENTOLGOVORETOPRNDELNFGKOAGECTES PQROLLKYSPERH JT COOTR	RPCTNRÇKACEPGESYBEVOPSYWRPPLMS RPCTNRÇKACEPGESYSEEVCROVPSYWRRPQMS RPCTNRQKACEPGESYSEEVCROVPSYWRRPQMS RPCTNRQKACEPGFSYSEEVCROVPSYWRRPQMS RPCTNRQKACEPGFSYSEEVCROVPSYWRRPQMS RPCTNRQKACEPGFSYSEEVCPSYWRPPQMS
F/L	F/2 (2) (3) (4)	(C) (C) (A) (A) (A) (A) (A) (A) (A) (A) (A) (A	(C) (C) (A)	(C) (C) (A) (A) (A) (A) (A) (A) (A) (A) (A) (A	$\frac{F}{(1)}$ (2) (3) (4)

Figure 2e pVORF1 MKLLVGILVAVCLHQYLLNADSNTKGWSEVIKGSE
LNADSNTKGWSEVILKGSE
NTKGWSEVILKGSE
SEVLKGSE
GCE

(3) (5) (5) (5) (5) (5)

CKERRIVVEVSETHRELITCORFNIPECVTLMROGGOCNPESLECVFTEEVNVTMELLGASGSGSNGMGRIGETURKTODO CKRRRIVVEVSETHPELITSQRFNPPCVTLMPOGGOCNPESLECVPTEEVNVTMELLGASGSGSNGMGRIGETURKTODO CKRRRIVVEVSETHPELITSQRFNPPCVTLMPOGGOCNPESLECVFTEEVNVTMELLGASGSGSNGMGRIGETURENGU CERRRIVYFTUSETHFELITSQRFNPPCVTLMPOGGOCNPESLECVFTEEVNVTMELLGASGSGSNGMGRIGETURENFROOG CKRPRIVVFVSRTHPELITSQRFNPPCVTLMROGGOCNPESLECVPTEEVNVTMELLGASGSGSNGMGRIGETURENFROOG	RPRETITEPRIRR RPRETITEPPRERR RPRETITEPITIRPPRER RPRETITEPITIRPPRERR
F/L	F/L
(1)	(2)
(3)	(3)
(4)	(4)

STNDWMRTLDKSG MRTLDKSG

MKLTATLQVVVALLICMYNLPECVSQSNDSPPSTNDWMRTLDKSG NDSPPSTNDWMRTLDKSG

F/L (2) (2) (3) (4)

Figure 2f pVORF2 CEPPTVVTLGEBYPESTYLQYIPPGVTVKFGSGOCHGEGOLGTAVETERITYTVSVTGVSSSSGTUSGVSTYLDEICVTERITRIDE ÖKPPDTVVYLGEBYPESTYLQYNPRGVTVKFGSGOCHGEGOLGTAVETRITTVTVSVTGVSSSSGTUSGVSTYLIGEISVTRHTRÄND ÖKPPDTVVYLGEBYPESTYLQYNPRGVTVKFGSGOCHGEGOLGTAVETRITTVTVSVTGVSSSSGTUSGVSTYLDEISVIEHTRÄDD ÖKPRDTVVYLGEBYPESTYLQYNPRGVTVKFGSGOCHGEGOLGTAVETRITTVTVSVTGVSSSGTUSGVSTYLQRISVIEHTKEND ÖKPRDTVVYLGEBYPESTYLQYNPRGVTVKFGSGOCHGEGOLGTAVETRITTVTVSVTGVSSSGTUSGVSTYLQRISVIEHTKEND

E/L (1) (2) (3) (3) (4)

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## WORLD INTELLECTUAL PROPERTY ORGANIZATION



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### (54) Title: TRUNCATED VEGF-RELATED PROTEINS

### VEGF-E

F/L (1) (2) (3) (4)	PVSQFDGPSHQKKVVPWIDVYTRAT PSHQKKVVPWIDVYTRAT KVVPWIDVYTRAT FWIDVYTRAT IDVYTRAT TDVYTRAT TTYRAT
(6)	RAT
F/L (1) (2) (3) (4) (5)	COPREVVVPLSMELMGNVVKOLVPSCVTVQRCGGCCPDDGLECVPTGCHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGCHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC CQPREVVVPLSMELMGNVVKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIQYPSSQLGEMSLEEHSQCEC
F/L (1)	RPKKKESAVKPOSPRILCPPCTQRRQRPDPRTCRCRCRRRRFLHCQGRGLELNPDTCRCRKPRK RPKKKESAVKPOSPRILCPPCTQRRQRPDPRTCRCRCRRRFFLHCQGRGLELNPDTCRCRKPRK
(2)	RPKKESAVKPDSPRILCPPCTORRORPDPRTCRCRCRRRFLHCOGRGLELNPDTCRCRKPRK
(3)	RPKKKESAVKPDSPRILCPPCTQRRQRPDPRTCRCRCRRRRFLHCQGRGLELNPDTCRCRKPRK
(4)	RPKFKESAVKPDSFRILCFPCTQRRQRPDPRTCRCRCRRRRFLHCQGRGLELNPDTCRCRKPRK
(5)	RPKFKESAVKPDSPRILCFPCTQRRQRPDPRTCRCRCRRRFLHCQGRGLELNPDTCRCRKPRK
(6)	RPKKKESAVKPDSPRILCPPCTQRRQRPDPRTCRCRCRRRRFLHCQGRGLELNPDTCRCRKPRK

### (57) Abstract

The present invention provides novel truncated forms of vascular endothelial growth factor-related proteins (VRPs or VRPs) which are useful for the stimulation of angiogenesis in vitro and in vivo. The invention also provides nucleic acids encoding such novel truncated VRPs and methods of producing truncated VRPs. Pharmaceutical compositions comprising truncated VRPs and methods of gene therapy using the nucleic acids which code for truncated VRPs may be useful for the treatment of heart disease and for wound healing.

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A. CLASSIFICATION OF SUBJECT MATTER IPC 6 012N15/12 0071 C07K14/52 C12N5/10 A61K38/19 According to international Patent Classification (iPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N C07K A61K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category 1 Relevant to claim No Citation of document, with indication, where appropriate, of the relevant passages χ WO 96 27007 A (AMRAD OPERATIONS PTY LTD 1-5. 17-23, (AU); HAYWARD NK; WEBER G; GRIMMOND S ET 25, AL.) 6 September 1996 28-30, 43-46. 48,49, 60,61 see abstract Υ 15,16, see page 6, line 24-25 32-36. 38, 40-42. 47, 50-59. 62-67 see page 10, line 17-21; figure 10 see page 19 - page 23; example 4 Seq.ID:4 see page 41 X Further documents are listed in the continuation of box  $\hat{C}$ [X ] Patent family members are listed in annex Special categories of cited documents To later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the last which is not cited to understand the principle or theory underlying the considered to be of particular resevance \*E\* earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to \*L\* document which may throw doubts on priority claim(s) or which is cred to establish the publication date of another involve an inventive step when the document is taken slone "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other, such docu-\*O\* document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means in the art. \*P\* document published prior to the international filling date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of making of the international search report a rie and mailing address of the ISM indized of tel Gropean Patent Office Fig. 1818 Patentiann. NL - 2280 HV Rijswick Tel. (+31-70) 340-2040, Tx 31 651 epo ni. Macchia, G Fax: (+31-70) 340-3016

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PCT/US 98/07801

	ation   DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
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,	see abstract see page 4, paragraph 6	15,16, 32-36, 38, 40-42, 47,56,59
	see page 6, paragraph 4 see page 10, paragraph 3 - page 12, paragraph 3 see page 18, paragraph 4 - page 20, paragraph 1 see page 22, paragraph 2 see page 25, paragraph 3 - page 27, paragraph 2 see page 38 - page 39; example 2 Seq.ID:2 see page 44	
, i	WO 96 26736 A (LUDWIG INST FOR CANCER RESEARCH (US); UNIV HELSINKI LICENSING LTD (FI)) 6 September 1996 see abstract see page 9, line 18-27 see page 10, line 29 - page 11, line 11 see page 18, line 3-26 see page 22 - page 23; example 4 see page 31 - page 33; example 9 see page 47, line 21 - page 48, line 30 see page 49, line 26 - page 50, line 5 Seq.ID:11,15 see page 59 - page 62 see figure 16	15,32, 33,56,59
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	(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT				
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Y	PEPPER M.S. ET AL.: "Potent synergism between vascular endothelial growth factor and basic fibroblast growth factor in the induction of angiogenesis in vitro" BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, vol. 189, no. 2, 15 December 1992, pages 824-831, XP002078851 see the whole document	50-55, 57,58			
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	see page 7, line 12-28 see page 8, line 28 - page 11, line 1 see page 15, line 11 - page 16, line 4 see page 27, line 14 - page 29, line 12 see page 30, line 21-31 see page 44 - page 45; example 7 see page 50, line 26 - page 53, line 15 see page 68 - page 74; examples 21,22 see page 84 - page 88; example 28 Seq.ID:33 see page 112 - page 113				
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	see page 6, paragraph 3 see page 21; claims 4,6 see page 24; claims 29,31				
Α	WO 94 11506 A (ARCH DEVELOPMENT CORPORATION; LEIDEN J.M.; BARR H. (US)) 26 May 1994	32-36, 38, 40-42, 47,62-67			
	see abstract	47,02 07			

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page : 11 3

Internation I application No

PCT/US 98/07801

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons
Claims Nos. because they relate to subject matter not required to be searched by this Authority, namely Remark: Although claim(s) 45, 47-52, 56-67 is(are) directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2. Claims Nos: because they relate to parts of the international Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically
Claims Nos because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6 4(a)
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows
See additional sheet
As all required additional search fees were time'y paid by the applicant, this International Search Report covers all searchable claims.
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As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos:
4 X No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims, it is covered by claims. Nos.
4 totally; 1-3, 5-67 partially.

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

1. Claims: 4 totally; 1-3, 5-67 all partially.

Truncated VEGF-B, VRF-2 or VEGF-3 subunit having a deletion of at least one of the aminoacid residues N-terminal to the first cysteine of the core sequence of said subunit, homodimers and heterodimers thereof. Nucleic acid molecule encoding said truncated subunit, recombinant vector comprising said nucleic acid molecule and transformed host cell thereof. Delivery vector comprising said nucleic acid molecule. Method for producing said truncated subunit. Pharmaceutical composition comprising said truncated subunit or delivery vector, therapeutical applications thereof.

2. Claims: 1-3, 5-67 all partially.

As invention 1 but concerning VEGF-C.

3. Claims: 1-3, 6-67 all partially.

As invention 1 but concerning PIGF.

4. Claims: 1-3, 5-67 all partially.

As invention 1 but concerning poxvirus ORF-1.

5. Claims: 1-3, 5-67 all partially.

As invention 1 but concerning poxvirus ORF-2.

Inform. In on patent family members

International Application No. PCT/US 98/07801

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